# OPPT NOIC

# 101-96-2 p-Phenylenediamine, N,N'-di-sec-Butyl

2.	PHYSICAL-CHEM	ICAL DATA	
*2.1	MELTING POINT		
	Value:	18°C	
	Decomposition:	Yes [] No [X] Ambiguous []	
	Sublimation:	Yes [ ] No [X] Ambiguous [ ]	
	Method:	Not determined	
	GLP:	Yes [ ] No [ ] ? [X]	
	Remarks:	HSDB, NTP Chemical Repository	
	Reference:	Ashford's Dictionary of Industrial Chemicals, 1994	
*2.2	BOILING POINT		
	Value:	98°C	
	Pressure:	at 26.6 hPa	
	Decomposition:	Yes [ ] No [X] Ambiguous [ ]	
	Method:	Not determined	
	GLP:	Yes [ ] No [ ] ? [X]	
	Remarks:	HSDB	
	Reference:	Kirk-Othmer Encyclopedia of Chemical Technology, 1991	
†2.3	†2.3 DENSITY (relative density)		
	Type:	Bulk density [X]; Density []; Relative Density []	
	Value:	0.94 kg/l	
	Temperature:	20°C	
	Method:	Not Determined	
	GLP:	Yes [ ] No [ ] ? [X]	
	Remarks:	HSDB	
	Reference:	Ashford's Dictionary of Industrial Chemicals, 1994	
*2.4	VAPOUR PRESSUE	RE	
	Value:	85.3 mm Hg	
	Temperature:	38°C	
	Method:	calculated [ ]; measured [X]	
		Instrumental method	
	GLP:	Yes [ ] No [ ] ? [X]	
	Remarks:	Radian Research	
	Reference:	NTP Chemical Repository, 2001	
*2.5	PARTITION COEF	FICIENT log <sub>10</sub> P <sub>ow</sub>	
	Log Pow:	3.50	
	Temperature:	Not determined	
	Method:	calculated [X]; measured []	
		SRC LogKow (KowWin) Program 1995	
	GLP:	Yes [ ] No [ ] ? [X]	
	Remarks:		
	Reference:	Meylan, W.M. and. P.H. Howard, 1995 J. Pharm. Sci. 84: 83-92	

# \*2.6 WATER SOLUBILITY

A.	Solubility			
	Value:	<1 mg/ml		
	Temperature:	20°C		
	Description:	Miscible []; Of very high solubility [];		
		Of high solubility []; Soluble []; Slightly soluble [];		
		Of low solubility []; Of very low solubility []; Not soluble [X]		
	Method:	Not determined		
	GLP:	Yes [ ] No [ ] ? [X]		
	Remarks:	Radian Research		
	Reference:	NTP Chemical Repository, 2001		
В.	pH Value, pKa Value			
	pH Value:			
	Concentration:			
	Temperature:	°C		
	Method:.			
	GLP:	Yes [] No [] ? []		
	pKa value	at 25°C		
	Remarks:			
	Reference:			
2.11	OXIDISING PROI	DEDTIES		
2.11	Results:	Maximum burning rate equal or higher than reference mixture[];		
	Results.	Vigorous reaction in preliminary test [ ];		
	Mathad.	No oxidising properties [ ]; Other [ ]		
	Method:	V [] N [] 0[]		
	GLP:	Yes [] No [] ? []		
	Remarks:			
	Reference:			
†2.12	OXIDATION: REI	DUCTION POTENTIAL		
	Value:	mV		
	Method:			
	GLP:	Yes [] No [] ? []		
	Remarks:			
	Reference:			
2.13	ADDITIONAL DA	TA		
<b>A.</b>	Partition co-efficien	t between soil/sediment and water (Kd)		
	Value:			
	Method:			
	GLP:	Yes [] No [] ? []		
	Remarks:			
	Reference:			
D	Other data			
В.				
	Results: Remarks:			
	Remarks:			

# 3. <u>ENVIRONMENTAL FATE AND PATHWAYS</u>

*3.1.1	PHOTODEGRADA	ATION				
	Type:	Air [ <b>X</b> ]; Water [ ]; Soil [ ]; Other [ ]				
	Light source:	Sunlight [ ]; Xenon lamp [ ]; Other [ ]				
	C I	nm				
		(based on intensity of sunlight)				
		ce: nm				
		ostance:				
	Temperature:	°C				
	Direct photolysis:					
	Half life:					
	Degradation:	% (weight/weight) after (exposure time)				
	Quantum yield:					
	Indirect Photolysis:					
	Type of sensitizer:					
		nsitizer: . 1560000 molecule/cm <sup>3</sup>				
		cal): 117.2377 E-12 cm <sup>3</sup> / molecule *sec				
	Degradation:	50% at 1.095 Hrs				
	Method:	calculated [X]; AOP Program (v1.89)				
		measured [ ]				
	GLP:	Yes [] No [X] ? []				
	Test substance:	. molecular structure, purity:				
	Remarks:	. morecular structure, parity				
	Rendomity. (2)	Accepted calculation method				
	Reference:	Meylan W. and Howard P. (1999) EPIWin Modeling Program.				
		Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.				
*3.1.2	STABILITY IN W.					
	Type:	Abiotic (hydrolysis) [ ]; biotic (sediment)[ ]				
	Half life:	at pH at°C				
	Degradation:	at pH at °C after				
		(exposure time)				
	Method:					
	GLP:	Yes [ ] No [ ] ? [ ]				
	Test substance:	purity:				
	Remarks:					
	Reference:					
*3.2	MONITORING DA	ATA (ENVIRONMENTAL)				
	Type of Measuremer	nt: Background [ ]; At contaminated site [ ]; Other [ ]				
	Media:					
	Results:					
	Remarks:					
	Reference:					
3.3	TRANSPORT A	ND DISTRIBUTION BETWEEN ENVIRONMENTAL				
	COMPARTMENT					
	CONCENTRATIO	NS AND DISTRIBUTION				

#### \*3.3.1 TRANSPORT

Type: Adsorption []; Desorption []; Volatility []; Other []

Media: Method: Results: Remarks: Reference:

# \*3.3.2 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION)

Media: Air-biota []; Air-biota-sediment-soil-water []; Soil-biota [];

Water-air []; Water-biota []; Water-soil []; Other []

Method: Fugacity level I [ ]; Fugacity level II [ ]; Fugacity level III [X];

Fugacity level IV[];Other (calculation)[];Other (measurement)[]

Results:

	Concentration	Half-Life	Emissions	Fugacity
	(percent)	(hr)	(kg/hr)	(atm)
Air	0.0952	2.19	1000	2.37e-012
Water	26.1	900	1000	2.36e-013
Soil	72.6	900	1000	2.33e-013
Sedime	ent 1.24	3.6e+003	0	1.75e-013

	Reaction	Advection	Reaction	Advection
	(kg/hr)	(kg/hr)	(percent)	(percent)
Air	678	21.4	22.6	0.714
Water	451	586	15	19.5
Soil	1.26e+003	0	41.9	0
Sediment	5.35	0.556	0.178	0.0185

Persistence Time: 750 hr Reaction Time: 940 hr Advection Time: 3.7e+003 hr

Percent Reacted: 79.7 Percent Advected: 20.3

Remarks:

Reliability: (2) valid with restrictions

Accepted calculation method

Reference: Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

# \*3.5 BIODEGRADATION

Type: aerobic [ ]; anaerobic [ ] Inoculum: adapted [ ]; non-adapted [ ];

Concentration of the chemical: . . . . . related to COD [ ]; DOC [ ]; test substance [ ] Medium: water [ ]; water-sediment [ ]; soil [ ]; sewage treatment [ ]

Degradation: (percentage reduction/exposure time)

..... % after ..... (time)

Results: readily biodeg. [ ]; inherently biodeg. [ ]; under test condition

no biodegradation observed [ ], other [ ]

Kinetic ...... % in ..... (time)

Method:

GLP: Yes [ ] No [ ] ? [ ]

Test substance: ...., purity: .....

Remarks: Reference:

# 4. <u>ECOTOXICITY</u>

# \*4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [X]

Species: <u>Salmo gairdneri</u> (Rainbow Trout)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = >0.18 mg/l

 $LC_{50}$  (48h) = 0.14 mg/l  $LC_{50}$  (72h) = Not determined  $LC_{50}$  (96h) = 0.13 mg/l NOEC = 0.056 mg/l LOEC = 0.10 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance:. Santoflex 44 dark red liquid #KB12-902, purity: >97%

Remarks: Acetone used as solvent. Range-finding experiment conducted to

determine final bioassay test concentrations. Water temperature, dissolved oxygen content, and pH monitored throughout study. Quality check via challenge with reference compound Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto AB-83X-036, Analytical Bio-Chemistry Labs, 1983

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [X]

Species: Lepomis machrochirus (Bluegill Sunfish)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 0.19 mg/l

 $LC_{50}$  (48h) = 0.18 mg/l  $LC_{50}$  (72h) = Not determined  $LC_{50}$  (96h) = 0.18 mg/l NOEC = 0.10 mg/l LOEC = 0.18 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance:. Santoflex 44 dark red liquid #KB12-902, purity: >97%

Remarks: Acetone used as solvent. Range-finding experiment conducted to

determine final bioassay test concentrations. Water temperature, dissolved oxygen content, and pH monitored throughout study. Quality check via challenge with reference compound Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto AB-83X-035, Analytical Bio-Chemistry Labs, 1983

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [X]

Species: Pimephales promelas (Fathead Minnows)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 0.13 mg/l

 $LC_{50}$  (48h) = 0.13 mg/l  $LC_{50}$  (72h) = Not determined  $LC_{50}$  (96h) = 0.13 mg/l NOEC = 0.10 mg/l LOEC = 0.18 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance:. Santoflex 44 dark red liquid #KB12-902, purity: >97%

Remarks: Acetone used as solvent. Range-finding experiment conducted to

determine final bioassay test concentrations. Water temperature, dissolved oxygen content, and pH monitored throughout study. Quality check via challenge with reference compound Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto AB-84X-021, Analytical Bio-Chemistry Labs, 1983

# 4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

# \*A. Daphnia

Type of test: static [X]; semi-static []; flow-through []; other [];

open-system [ ]; closed-system [X]

Species: <u>Daphnia magna</u>

Exposure period: 48 Hours

Results:  $EC_{50}$  (24h) = 2.0 mg/l

 $EC_{50}$  (48h) = 1.4 mg/l NOEC = 0.56 mg/l

Analytical monitoring: Yes [X] No [ ] ? []

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex 44 dark liquid Lot# KB12-902, purity:>97%

Remarks: Acetone used as solvent. Range-finding experiment conducted to

determine final bioassay test concentrations. Water temperature, dissolved oxygen content, and pH monitored throughout study. The abnormal effects of mortality and daphnids lying on the bottom progressed from 3.2 mg/l initially, to 1.0 mg/l after 48

hours.

Reference: Monsanto AB-83X-037, Analytical Bio-Chemistry Labs, 1983

# \*4.3 TOXICITY TO AQUATIC PLANTS, e.g. algae

Species:

Endpoint: Biomass []; Growth rate []; Other []

Exposure period:

Results:  $EC_{50} (\dots h) = \dots mg/l$ 

 $EC_{xx}$  (......h) = ......mg/l

 $NOEC = \dots mg/l$  $LOEC = \dots mg/l$ 

Analytical monitoring: Yes [ ] No [ ] ? [ ]

Method:

open-system []; closed-system []

GLP: Yes [ ] No [ ] ? [ ]

Test substance: ...., purity:....

Remarks: Reference:

# 5. TOXICITY

#### \*5.1 ACUTE TOXICITY

#### 5.1.1 ACUTE ORAL TOXICITY

Type:  $LD_0[]$ ;  $LD_{100}[]$ ;  $LD_{50}[X]$ ;  $LDL_0[]$ ; Other[]

Species/strain: Sprague-Dawley Albino Rats

Value: 271 mg/kg bw for males and females combined

281 mg/kg for males 265 mg/kg for females

Method: Finney, J.D., Reference for Method of LD50 Determination,

Probit Analysis 3rd Edition, 1971

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 44, Lot S-40182, purity: 96.09%

Remarks: Groups of five male and five female rats were dosed by oral

gavage with the test article as a 392 mg/ml solution in corn oil. Clinical observations were made 3x/day during the first 8 hours, and 2x/day thereafter until sacrifice. After a 14-day recovery period, all surviving animals were sacrificed. Necropsies were performed on all animals. Clinical signs of toxicity included lethargy, ataxia, ptosis, and abnormal urine coloration (green reddish-brown). Necropsy findings and/or included gastrointestinal inflammation, which reached the severity of hemmorhage in many cases, gastrointestinal distension, and red, fluid-filled gastric masses. The presence of these masses indicated that the toxicity to gastrointestinal tissue may have contributed to lethality in virtually all rats that died during the test. Previous oral and dermal toxicity studies with this material have noted the corrosivity to tissue that complicates accurate

determinations of LD50 values.

Reference: Monsanto ML-82-181, Environmental Health Labs, 1983

#### 5.1.2 ACUTE INHALATION TOXICITY

Type:  $LC_0[]; LC_{100}[]; LC_{50}[]; LCL_0[X]; Other[]$ 

Species/strain: Sprague-Dawley Albino Rats

Exposure time: 6 Hours
Value: 600 mg/m3
Method: Not Determined

GLP: Yes [ ] No [ ] ? [ X ] Klimisch 2
Test substance:. Cas # 101-96-2, purity: Commercial (>96%)

Remarks: RTECS and NTP reference. Test conditions unknown.

Reference: Kodak Company Reports, 1971

Type:  $LC_0[]; LC_{100}[]; LC_{50}[X]; LCL_0[]; Other[]$ 

Species/strain: Sprague-Dawley Albino Male Rats

Exposure time: 6 Hours
Value: >0.2 mg/l
Method: A.T.S. 8/1973

GLP: Yes [ ] No [X ] ? [ ] Klimisch 2
Test substance:. Santoflex 44 Lot# 24277, purity: >96%

Remarks: Six male rats were exposed to the test article at a concentration

of 0.2 mg/l at ambient temperature at an airflow rate of 4 l/min for six hours. The difference in weight of the sample after the test indicated that 0.4 grams had been vaporized under test conditions. There were no clinical signs of toxicity noted during the experiment. Following a 14-day recovery period, all animals were sacrificed. Necropsy findings were that all viscera examined

appeared normal.

Reference: Monsanto Y-76-262, Younger Laboratories, 1976

#### 5.1.3 ACUTE DERMAL TOXICITY

Type:  $LD_0[]; LD_{100}[]; LD_{50}[X]; LDL_0[]; Other[]$ 

Species/strain: New Zealand Albino Rabbits

Value: 2806 mg/kg bw

Method: Finney, J.D., Reference for Method of LD50 Determination,

Probit Analysis 3<sup>rd</sup> Edition, 1971

GLP: Yes [X] No []?[] Klimisch 1
Test substance: Santoflex 44 Lot S-40182, purity: 96.09%

Remarks: Groups of four male and female rabbits were exposed to the test

article via a single dermal application to shaved skin. Two animals from each group were predesignated to have their skin abraided in the treatment area. Skin of the other animals was intact. Clinical observations were made 3x/day during the first eight hours after exposure, then 2x/day thereafter until sacrifice. Necropsies were performed on all animals. Clinical signs of toxicity included lethargy, ataxia, green coloration of the urine, partial loss of ability to move the limbs, and localized dermal effects attributed to the direct contact between skin and test article. Findings on necropsy included green material in the bladder of sixteen animals, four animals with an enlarged gall

bladder, and five with hepatic discoloration.

Reference: Monsanto ML-82-022, Environmental Health Lab, 1983

# 5.2 CORROSIVENESS/IRRITATION

#### 5.2.1 SKIN IRRITATION/CORROSION

Species/strain: New Zealand White Rabbits

Results: Highly corrosive [ ]; Corrosive [ X ]; Highly irritating [ ];

Irritating [ ]; Moderate irritating [ ]; Slightly irritating [x ];

Not irritating [ ]

Classification: Highly corrosive (causes severe burns) [ ];

Corrosive (causes burns) [X]; Irritating []; Not irritating []

Method: Draize, J.H. Woodard, G., and Calvery, H.O., Methods for the

Study of Irritation and Toxicity of Substances Applied Topically

To the Skin and Mucous Membranes, J. Pharmacol. Exp.

Therap. 82: 377-390, 1944

GLP: Yes [X] No [ ] ? [ ] Klimisch 1
Test substance: Santoflex 44 Lot S-40182, purity: 96.09%

Remarks: The test undiluted article, at a volume of 0.5 ml, was applied to

the intact and abraided shaved skin of six rabbits for 24 hours. The initial observation was made approximately one hour after exposure. Dermal irritation was scored by the Draize Method, and results recorded on day 1, 3, 7, 10, 14 and 17 after exposure. Scarring, hardening of the skin, scabbing and sloughing skin were noted on all animals. The test article was classified as

corrossive under the test conditions.

Reference: Monsanto ML-82-022c, Environmental Health Lab, 1983

#### SKIN IRRITATION/CORROSION

Species/strain: New Zealand Albino Rabbits

Results: Highly corrosive [ ]; Corrosive [ ]; Highly irritating [ X ];

Irritating [ ]; Moderate irritating [ ]; Slightly irritating [ ];

Not irritating [ ]

Classification: Highly corrosive (causes severe burns) [ ];

Corrosive (causes burns) [ ]; Irritating [X]; Not irritating [ ]

Method: D.O.T. Hazardous Material Regulations 49 CFR 173.240, 1976

GLP: Yes [ X ] No [ ] ? [ ] Klimisch 1

Test substance:. Antioxidant PDA #1549-83, purity: Not stated

Remarks: The undiluted test article was applied to the shaved skin of six

rabbits in a single application of 0.5 ml. The test site was covered for four hours with surgical gauze and an elastic bandage. The entire trunk of the rabbit was wrapped in 2 mil thick plastic to prevent evaporation of the test article, and the plastic was covered with a white cotton towel. After four hours, the wrappings were removed, and the skin allowed to equilibrate for hydration and compression for 30 minutes. Skin was scored for erythema, eschar formation and corrosion in accordance with the Federal Hazardous Substances Act Grading Code, 16 CFR 1500.41. After grading, the test site was washed with water. Test sites were scored again after 24, 48 and 72 hours, and 1 and 2 weeks. Gross observations of corrosion were noted in 2/6 rabbits at I week and in 4/6 rabbits after 2 weeks. Under the conditions of the DOT test, these results were judged to be between "marginal" and "severely irritating but not corrosive". Because of the results of earlier studies, the manufacturers of this material have chosen to classify it as "corrosive" for both use and

transportation.

Reference: Monsanto XX-84X-144, Gulf South Research, 1983

#### 5.2.2 EYE IRRITATION/CORROSION

Species/strain: New Zealand Albino Rabbits

Results: Highly corrosive [ ]; Corrosive [ X ]; Highly irritating [ ];

Irritating [ ]; Moderate irritating [ ]; Slightly irritating [ ];

Not irritating [ ]

Classification: Irritating []; Not irritating []; Risk of serious damage to eyes [X] Method: Draize et.al., <u>J. Pharmacol., Exp. Therap.</u> 82: pp 377-390, 1944

GLP: Yes [] No [] ? [x]

Test substance:. Santoflex 44 Lot# S-40182, purity 96.09%

Remarks: A single dose of 0.1 ml of the undiluted test article was placed in

the one eye of three male and three female rabbits, with the untreated eye serving as the control. A topical anesthetic available if discomfort appeared severe. Signs of irritation were scored according to the Draize procedure. Scoring will be done at 24, 48 and 72 hours after treatment. Discomfort on application was slight. Observations at 24 hours included severe erythema with necrosis, severe edema, copious discharge containing a whitish exudate and severe swelling of conjunctivae. Under the test conditions, the material was classified as "corrosive". Scabs sloughed off in 14 to 21 days with no apparent permanent

corneal damage.

Reference: Monsanto ML-82-022d, Environmental Health Laboratory, 1983

# \*5.4 REPEATED DOSE TOXICITY

Species/strain: Sprague-Dawley Albino Rats

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral gavage

Exposure period: 28 days
Frequency of treatment: Daily
Post exposure observation period:

Dose: 0, 10, 25, 50, or 100 mg/kg
Control group: Yes [ **X** ]; No [ ]; No data [ ];

Concurrent no treatment []; Concurrent vehicle [X]; Historical []

NOEL: <10 mg/kg LOEL: 10 mg/kg

Results: 100 male and female rats (10/sex/dose level) were dosed with the

test article in corn oil vehicle at the above levels for a period of 28 days. The animals were observed 2x/day for mortality or signs of toxicity. Detailed observations, body weights and feed consumption was documented 1x/week. Hematology determinations and clinical chemistry determinations were made on all control animals and the high-dose animals prior to terminal sacrifice. Additional clinical chemistry determinations of GGTP, SGOT, Sgtp, Bilirubin, SAP and 5-nucleotidase were performed on all treated animals. A complete gross necropsy was performed on all animals at sacrifice and within 16 hours of any animal who died during the course of the study. Two mid-dose males died within the first week of treatment and two high-dose females died during week 3. Cause of death did not appear to be treatmentrelated. One additional mid-dose female was sacrificed at day 15 following an injury during dosing. All other animals survived to sacrifice. Gross necropsy findings on two high-dose females was a slightly pale liver. In males, a finding of dilation of the right renal pelvis was found in several animals at all dose levels, including controls. Adverse effects observed included increased liver weights and elevation of serum enzymes SGOT, Sgpt and GGTP, indicative of hepatocellular damage, as well as a dosedependent increase in the incidence of hepatocellular lesions. Because the results of this study demonstrated hepatic effects in

both sexes and at all treatment levels, a No Observed Effect

Level could not be established.

Method: OECD Guidelines for the Testing of Chemicals, 1981

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 44 Lot# KC11-928, purity: >96%

Reference: Monsanto PR-83-317, Pharmacopathics Research Labs, 1984

Species/strain: Sprague-Dawley Albino Rats

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral dietary

Exposure period: 90-94 days Frequency of treatment: Daily Post exposure observation period:

Dose: 0, 20, 100 or 500 ppm

Control group: Yes [X]; No []; No data [];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL: 100 mg/kg LOEL: 500 mg/kg

Results: In a subchronic feeding study, groups of male and female rats

were fed the test article via dietary admixture for three months. After 65 days of treatment, the low-dose (20 ppm) group was increased to 1000 ppm for twenty-five days, and then to 2000 ppm for the final four days of the study. Findings included decreased body weights and body weight gain in the 500

ppm males, and decreased body weights in the 500 ppm females. There were no clinical signs of toxicity noted for any dose level for either sex. All animals survived until terminal sacrifice. Hematology determinations and clinical chemistry determinations were made on all animals prior to sacrifice, and all animals

received a complete gross necropsy. There were no

hematological or histopathological findings at any dose level that were considered to be treatment-related. The NOEL was determined to be 100 ppm, or 6.6 mg/kg/day, for both males and females based upon the reduced body weights seen at 500 ppm.

Method: Not determined

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: Antioxidant 22, purity: Commercial grade, 96% minimum

Reference: E.I. DuPont de Nemours, unpublished data, 1987

# \*5.5 GENETIC TOXICITY IN VITRO

#### A. BACTERIAL TEST

Type: Bacterial Reverse Mutation - Ames

System of testing: Salmonella typhimurium TA97, TA98, TA100, TA1535, TA

1537, TA1538

Concentration: Not determined

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: Not determined

Without metabolic activation: Not determined

Precipitation conc: Not determined Genotoxic effects: + ? -With metabolic activation: [][][X] Without metabolic activation: [][][X] Method: OECD 471 Plate Overlay method GLP: Yes [] No [] ? [X] Klimisch 2 N,N'-di-sec-butyl-p-phenylenediamine, purity: Technical grade Test substance: The test compound was tested in Ames/Salmonella plate Remarks: incorporation assays using the tester strains TA 97, TA98, T A100, TA1535, and TA1538 and TA1537 in the presence and absence of an Aroclor-induced rat liver mammalian metabolic activation system (S-9 Mix). No mutagenic activity was observed for the test compound in any of these assays. Reference: Zeiger, et. al., Environ. Mol. Mutagen, 1998 В. NON-BACTERIAL IN VITRO TEST CHO and CHL Forward Gene Mutation Assay Type: System of testing: Cultured Chinese hamster ovary (CHO) cells and cultured Chinese Hamster Lung (CHL) cells Concentration: Not determined Metabolic activation: With []; Without []; With and Without [X]; No data [] Results: Cytotoxicity conc: With metabolic activation: Not determined Without metabolic activation: Not determined Precipitation conc: Not determined Genotoxic effects: With metabolic activation (CHO): [][][X] Without metabolic activation (CHO): [][][X] With metabolic activation (CHL): [][][X] [][X][] Without metabolic activation (CHL): Method: **OECD 476** GLP: Yes [ ] No [ ] ? [X] Klimisch 2 N,N'-di-sec-butyl-p-phenylenediamine, purity: Commercial grade Test substance:. Remarks: The test article was one of 25 chemicals tested for the induction of chromosomal aberrations in two cultured mammalian cell systems the cultured cells from Chinese hamster ovaries (CHO, and those from Chinese hamster lungs (CHL), in the presence absence of metabolic activation with the S9 mix. The test article negative with metabolic activation in both CHO and CHL cells, and negative without metabolic activation in CHO cells. The results for CHL cells without metabolic activation were equivocal. Overall, the results indicate that the test article is negative for the potential to cause chromosomal aberrations, both with and without metabolic activation, under the test conditions. Reference: Sofuni, et.al. Mutation Research, 1990 \* 5.6 GENETIC TOXICITY IN VIVO

Type:

Species/strain:

Sex: Female []; Male []; Male/Female []; No data []

Route of Administration:

Exposure period:

	Doses.			
	Results:			
	Effect on mitotic			
	index or P/N ratio:			
	Genotoxic effects:			
		[][][]		
	Method:			
	GLP:	Yes [ ] No [ ] ? [ ]		
	Test substance:	, purity:		
		, purity		
	Remarks:			
	Reference:			
*5.8	TOXICITY TO REP	PRODUCTION		
	Type:	Fertility []; One-generation study []; Two-generation study [];		
	· -	Termity [], one generation study [], 1 wo generation study [],		
	Species/strain:			
	Sex:	Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]		
	Route of Administration	on:		
	Exposure period:			
	Frequency of treatmen	f•		
	Post exposure observat			
	*	*		
		eriod: male: , female:		
	Duration of the test:			
	Doses:			
	Control group:	Yes [ ]; No [ ]; No data [ ];		
	8 11	Concurrent no treatment [ ]; Concurrent vehicle [ ]; Historical [		
1	NOEL Parantal	Concurrent no treatment [ ], Concurrent venicle [ ], Instoricar [		
]	NOEL Parental:	•		
	NOEL F1 Offspring:			
	NOEL F2 Offspring:			
	Results:			
		General parental toxicity:		
		Toxicity to offspring:		
	Mathad.	Toxicity to offspring.		
	Method:			
	GLP:	Yes [ ] No [ ] ? [ ]		
	Test substance:	, purity:		
	Remarks:			
	Reference:			
*5.9	DEVELOPMENTAL TOXICITY/ TERATOGENICITY			
	Species/strain:			
	Sex:	Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]		
	Route of Administration	on: .		
	Duration of the test:			
	Exposure period:			
	Frequency of treatmen	<b>f•</b>		
	ž •	L.		
	Doses:			
	Control group:	Yes [ ]; No [ ]; No data [ ];		
	- <b>-</b>	Concurrent no treatment []; Concurrent vehicle []; Historical []		
	NOEL Maternal Toxic			
	NOEL teratogenicity:			
	Results:			
		Maternal general toxicity:		
		Pregnancy/litter data:		

]

Foetal data:

Method:

GLP: Yes [ ] No [ ] ? [ ]

Test substance: ...., purity: .....

Remarks: Reference:

# 5.10 OTHER RELEVANT INFORMATION

#### A. Specific toxicities

Type: Results: Remarks: Reference:

# B. Toxicodynamics, toxicokinetics

Type: Results: Remarks: References:

#### \* 5.11 EXPERIENCE WITH HUMAN EXPOSURE

Results: Cyanosis and anemia have been observed in workers involved in the

manufacture of Antioxidant 22.

Remarks: Dermal route

Reference: E,I, DuPont de Nemours, 1987

Results: Historically, three incidents involving accidental human overexposure

involving Antioxidant 22 have been documented. Skin reactions noted were irritation and a pigmented crust that scaled away in a few days, leaving an erythematous base. Systemic reactions, indicative of skin absorbtion, included profuse perspiration, slow pulse, and a general

feeling of anxiety.

Remarks: Data from 1945 does not reflect current industrial practice utilizing

Impervious gloves and other personal protective equipment

Reference: Kendrick, M.C., The Medical Bulletin, 1945

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# 3081-14-9

# p-Phenylenediamine, N-1,4-Dimethylpentyl-N'-Phenyl-

# 2. PHYSICAL-CHEMICAL DATA

\*2.1 MELTING POINT

Value: -36 °C

Decomposition: Yes [ ] No [X ] Ambiguous [ ] Sublimation: Yes [ ] No [X ] Ambiguous [ ]

Method: Not Specified

GLP: Yes [ ] No [ ] ? [X]

Remarks:

Reference: NTP Chemical Repository 1990

\*2.2 BOILING POINT

Value: 183 °C Pressure: 1mm Hg

Decomposition: Yes [ ] No [X] Ambiguous [ ] Method: Capillary Melt-Temp Instrument

GLP: Yes [ ] No [ ] ? [X]

Remarks:

Reference: Monsanto Physical Constants of CP25447 (SMP 1977)

†2.3 DENSITY (relative density)

Type: Bulk density []; Density [X]; Relative Density []

Value: 0.9 Temperature: 27 °C

Method: Flexsys Standard Method of Analysis FF97.4-1

GLP: Yes [ ] No [ ] ? [X]

Remarks: Hydrometer method. Hydrometer must meet standards set in

ASTM-E-100

Reference: ASTM D891-94 method equivalent

\*2.4 VAPOUR PRESSURE

Value: <1.1 x 10(-6) Torr

Temperature: 25°C

Method: calculated [ ]; measured [X]

Gas Saturation Method, W.F. Spencer and M.M. Cliath, Environ. Sci.

Tech. <u>3</u>, 670 (1969)

GLP: Yes [X] No [] ? []

Remarks: Nitrogen carrier gas, Tenax-GC sorbent, GC analysis

Reference: Monsanto SRI 8669, SRI International, 1980

\*2.5 PARTITION COEFFICIENT log<sub>10</sub>P<sub>ow</sub>

Log Pow: 5.34 log P
Temperature: 22°C

Method: calculated []; measured [X]

EPA Federal Register Vol. <u>44</u>, No. 53 (1979)

GLP: Yes [X] No [] ? [] Remarks: Octanol used as solvent

Reference: Monsanto SRI 8669, SRI International, 1980

# \*2.6 WATER SOLUBILITY

A. Solubility

Value: 21 ppm @ pH 5, 0.8 ppm @ pH 9

Temperature: 22°C

Description: Miscible []; Of very high solubility [];

Of high solubility []; Soluble []; Slightly soluble [];

Of low solubility []; Of very low solubility [X]; Not soluble []

Method: May, W.E., Wasik, S.P., Freeman, D.H., Anal. Chem. <u>50</u> (1)

175-178, 1978

GLP: Yes[X] No[]?[]

Remarks: May Method chosen for low-solubility chemicals Reference: Monsanto SRI 8669, SRI International, 1980

B. pH Value, pKa Value

pH Value: Not Applicable

**2.7 FLASH POINT** (liquids)

Value: 182 °C

Type of test: Closed cup []; Open cup [X]; Other []
Method: ASTM D 92 Cleveland Open Cup

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Remarks: No method deviations

Reference: American Society for Testing and Materials, 1997

2.13 ADDITIONAL DATA

A. Partition co-efficient between soil/sediment and water (Kd)

Value: Method:

GLP: Yes [ ] No [ ] ? [ ]

Remarks: Reference:

3. <u>ENVIRONMENTAL FATE AND PATHWAYS</u>

3.1 STABILITY

\*3.1.1 PHOTODEGRADATION

Type: Air [ ]; Water [ X ]; Soil [ ]; Other [ ]
Light source: Sunlight [X]; Xenon lamp [ ]; Other [ ]

Light spectrum: Natural sunlight, March 7, 1980

Relative intensity:

Spectrum of substance: 262 nm Concentration of Substance: 5ppm Temperature: 23 °C

Direct photolysis:

Half life: 2 hours (light) and 4 hours (dark)

Degradation: Quantum yield:

Method: calculated [ ]; measured [X]

**Direct Photolysis** 

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex 77 dark liquid, purity: >94%

Remarks:

Reference: Monsanto SR-85-017 SRI International, 1985

Type: Air [X]; Water [ ]; Soil [ ]; Other [ ]
Light source: Sunlight [ ]; Xenon lamp [ ]; Other [ ]

Light spectrum: .....nm

Relative intensity: ..... (based on intensity of sunlight)

Direct photolysis:

Half life: .....

Degradation: ..... % (weight/weight) after ..... (exposure time)

Quantum yield: .....

Indirect Photolysis:

Type of sensitizer: .....OH ...

Concentration of sensitizer: . . 1560000 . . molecule/. cm<sup>3</sup> . . . . . Rate constant (radical): . . . 125.6992 E-12. . . cm<sup>3</sup>/molecule\*sec

Degradation: ... 50% at 1.021 Hrs. ....

Method: calculated [X]; AOP Program (v1.89)

measured [ ]

GLP: Yes [] No [X]? []

Test substance: ......, purity:..... purity:.....

Remarks:

Reliability: (2) valid with restrictions

Accepted calculation method

Reference: Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

#### \*3.1.2 STABILITY IN WATER

Type: Abiotic (hydrolysis) [ X ]; biotic (sediment)[ ]

Half life: Not measured

Degradation: 97% at pH 7.0 at 25 °C after 24 hours exposure time Method: Phase I Hydrolysis Study / ID of Hydrolysis Products

GLP: Yes [X] No [] ? [] Klimisch 1
Test substance: Santoflex 77 dark reddish liquid, purity: >94%

Remarks: Rapid hydrolysis to 4-Hydroxylamine and Benzoquinoneimine-N-phenyl.

No test substance detected after 7 days.

Reference: Monsanto ABC-32303 Analytical BioChemistry Labs 1986

# \*3.3.2 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION)

Media: Air-biota []; Air-biota-sediment-soil-water []; Soil-biota [];

Water-air []; Water-biota []; Water-soil []; Other []

Method: Fugacity level I [ ]; Fugacity level II [ ]; Fugacity level III [ X]; Fugacity

level IV [ ]; Other (calculation) [ ]; Other (measurement)[ ]

Results:

	Concentration	Half-Life	Emissions	Fugacity
	(percent)	(hr)	(kg/hr)	(atm)
Air	0.0609	2.04	1000	2.1e-012
Water	5.53	900	1000	1.65e-013
Soil	31.7	900	1000	1.25e-015
Sediment	62.7	3.6e+003	0	1.11e-013

	Reaction	Advection	Reaction	Advection
	(kg/hr)	(kg/hr)	(percent)	(percent)
Air	901	26.6	30	0.885
Water	186	241	6.19	8.04
Soil	1.06e+003	0	35.5	0
Sediment	527	54.7	17.6	1.82

Persistence Time: 1.45e+003 hr Reaction Time: 1.63e+003 hr Advection Time: 1.35e+004 hr

Percent Reacted: 89.3 Percent Advected: 10.7

Remarks:

Reliability: (2) valid with restrictions

Accepted calculation method

Reference: Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

#### \*3.5 BIODEGRADATION

Type: aerobic [X]; anaerobic []

Inoculum: adapted [ X ]; non-adapted [ ]; Sewage/soil/sludge mixture

Concentration of the chemical: 25 mg/l related to COD [ ]; DOC [ ]; test substance [X]

Medium: water []; water-sediment []; soil []; sewage treatment [ X ]

Degradation: 50% of theory after 35 days

Results: readily biodeg. []; inherently biodeg. [X]; under test condition no

biodegradation observed [ ], other [ ]

Kinetic ...... % in ..... (time)

Method: ASTM Proposed Standard for the Determination of the Ultimate

Biodegradability of Organic Chemicals, 1979

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex 77 Lot# KL01-04, purity:>94%

Remarks: Sterile controls used – no significant biodegradation noted under sterile

conditions. Test run in triplicate.

Reference: Monsanto ES-79-SS-25 MIC Environmental Sciences, 1979

# 4. <u>ECOTOXICITY</u>

#### \*4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type of test: static [X]; semi-static []; flow-through []; other (e.g. field test) []

open-system [ ]; closed-system [X ]

Species: Salmo gairdneri (Rainbow Trout)

Exposure period: 96 hours

Results:  $LC_{50}$  (24h) = 51 mg/l

 $LC_{50}$  (48h) = 39 mg/l

 $LC_{50}$  (72h) = Not Measured

 $LC_{50}$  (96h) = 32 mg/l NOEC = 20 mg/l LOEC = 32 mg/l

Analytical monitoring: Yes [X] No [] ? []

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975)

GLP: Yes [ ] No [ ] ? [X ] Klimisch 1

Test substance:. Santoflex 77 dark red liquid Lot #KD05-57 purity:>94%

Remarks: Acetone used as solvent. Water temperature, dissolved oxygen content,

and pH monitored throughout study. Data reported at 95% confidence

level.

Reference: Monsanto BN-76-254 EG&G Bionomics, 1976

Type of test: static [X]; semi-static []; flow-through []; other (e.g. field test) []

open-system [ ]; closed-system [X ]

Species: <u>Lepomis machrochirus</u> (Bluegill Sunfish)

Exposure period: 96 hours

Results:  $LC_{50}$  (24h) = 261 mg/l

 $LC_{50}$  (48h) = 201 mg/l  $LC_{50}$  (72h) = Not Measured  $LC_{50}$  (96h) = 182 mg/l NOEC = 140 mg/l LOEC = 180 mg/l

Analytical monitoring: Yes [X] No [] ? []

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975)

GLP: Yes [ ] No [ ] ? [X ] Klimisch 1

Test substance:. Santoflex 77 dark red liquid Lot #KD05-57 purity:>94%

Remarks: Acetone used as solvent. Water temperature, dissolved oxygen content,

and pH monitored throughout study. Data reported at 95% confidence

level.

Reference: Monsanto BN-76-254 EG&G Bionomics, 1976

Type of test: static [X]; semi-static []; flow-through []; other (e.g. field test) []

open-system [ ]; closed-system [X ]

Species: <u>Pimephales promelas</u> (Fathead Minnows)

Exposure period: 96 hours

Results:  $LC_{50}$  (24h) = 0.32 mg/l

$$\begin{split} LC_{50} \ (48\text{h}) &= 0.28 \ mg/l \\ LC_{50} \ (72\text{h}) &= Not \ Measured \\ LC_{50} \ (96\text{h}) &= 0.28 \ mg/l \\ NOEC &= Not \ Determined \\ LOEC &= 0.10 \ mg/l \end{split}$$

Analytical monitoring: Yes [X] No [] ? []

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex 77 dark red liquid purity 99+%

Remarks: Acetone used as solvent. Water temperature, dissolved oxygen content,

and pH monitored throughout study. Data reported at 95% confidence level. Quality check via Antimycin A challenge. Preliminary 72-hour

range-finding study used to determine final concentrations.

Reference: Monsanto AB-79-1384361-1a, Analytical BioChemistry Labs, 1979

# 4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

\*A. Daphnia

Type of test: static [X]; semi-static []; flow-through []; other (e.g. field test) [];

open-system [ ]; closed-system [ X ]

Species: <u>Daphnia magna</u>

Exposure period: 48 hours

Results:  $EC_{50}$  (24h) = 0.44 mg/l

 $EC_{50}$  (48h) = 0.37 mg/l NOEC = 10 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1
Test substance: Santoflex 77 reddish-brown liquid, purity: 99+%

Remarks: Nanograde Acetone used to prepare stock solutions. Water quality

parameters (temperature, dissolved oxygen, pH) monitored throughout study. Initial range-finding experiment used to select concentrations. Data

reported at 95% confidence level.

Reference: Monsanto AB-79-1384361-1b Analytic Bio-Chemistry Labs, 1979

B. Other aquatic organisms

Type of test: static [X]; semi-static []; flow-through []; other (e.g. field test) [];

open-system [ ]; closed-system [X]

Species: Paratanytarsus parthenogenetica (Midge)

Exposure period: 48 hours

Results:  $EC_{50}$  (24h) = 4.4 mg/l

 $EC_{50}$  (48h) = 1.7 mg/l NOEC = 0.56 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1
Test substance:. Santoflex 77 dark liquid, purity:>94%

Remarks: Stock solutions prepared in acetone. Range-finding experiment run to

determine final experimental concentrations. Water quality parameters

monitored throughout testing.

Reference: Monsanto AB-81-9AB981014, Analytical BioChemistry Labs, 1981

\*4.3 TOXICITY TO AQUATIC PLANTS, e.g. algae

Species: <u>Selenastrum capricornutum</u> (Freshwater alga)
Endpoint: Biomass [ X ]; Growth rate [ ]; Other [ ]

Exposure period: 96 hours

Results:  $EC_{50}$  (24h) = >200 mg/l

 $EC_{50}$  (48h) = >120<200 mg/l

 $EC_{50}$  (72h) = 86 mg/l  $EC_{50}$  (96h) = 52 mg/l

> NOEC = Not Determined LOEC = Not Determined

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA <u>Selenastrum capricornutum</u> Algal Assay Test 1978

open-system []; closed-system [X]

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 77 blackish-red liquid, Lot# KL01-04, purity: 99+%

Remarks: Stock solutions prepared in DMSO. Both cell numbers and decrease of in

vivo chlorophyll a measured. Triplicate cultures employed for all test

concentrations and for controls. pH monitored throughout test.

Reference: Monsanto BN-79-1384361-2, EG&G Bionomics, 1979

# 4.5 CHRONIC TOXICITY TO AQUATIC ORGANISMS

**4.5.1 CHRONIC TOXICITY TO FISH** (effects on reproduction, embryo/larva, etc.)

Type of test: static []; semi-static []; flow-through [X]; other (e.g. field test) []; open-

system [ ]; closed-system [x]

Species: <u>Pimephales promelas</u> (Fathead Minnow) Endpoint: Length of fish [ ]; Weight of fish [X];

Reproduction rate [ ]; Other [ ]

Exposure period: 14 days

Results:  $EC_{50}$  (14d) = 0.067 mg/l

NOEC = 0.018 mg/l LOEC = 0.046 mg/l

Analytical monitoring: Yes [X] No []?[]

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975) and Committee on Methods for Toxicity Tests with

Aquatic Organisms, 1975

GLP: Yes [X] No []? [] Klimisch 1

Test substance: Santoflex 77 dark liquid, purity: 99+%

Remarks: Stock solutions prepared in Methanol. Water quality parameters monitored

throughout test and remained withing acceptable limits. Behavior observations throughout the test indicated that mortality was preceded by surfacing and loss of equilibrium. Weight measurements of surviving fish at the end of the study yielded the following weight percentages of the control group mean weight: 0.018 mg/l = 84%, and 0.046 mg/l = 81%. An apparent lethal threshold of the test substance to fathead minnows was determined to be 0.067 mg/l and was reached after 12 days as indicated by

a cessation in mortality from days 12-14.

Reference: Monsanto AB-80-1803058-B1, Analytical BioChemistry Labs, 1981

Type of test: static []; semi-static []; flow-through [X]; other (e.g. field test) []; open-

system [ ]; closed-system [x]

Species: <u>Pimephales promelas</u> (Fathead Minnow)
Endpoint: Length of fish [X]; Weight of fish [X];

Reproduction rate [ ]; Other [ ]

Exposure period: 14 days (336 hours) Results:  $LC_{50}$  (24h) = 0.07 mg/l  $LC_{50}$  (96h) = 0.06 mg/l  $LC_{50}$  (14d) = 0.05 mg/l NOEC = Not DeterminedLOEC = Not Determined

Analytical monitoring: Yes [X] No []?[]

Method: EPA Methods for Acute Toxicity Tests with Fish, Macroinvertebrates and

Amphibians (1975) and Committee on Methods for Toxicity Tests with

Aquatic Organisms, 1975

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 77 dark reddish liquid, purity: 99+%

Remarks: Stock solutions prepared in acetone and stabilized with ascorbic acid.

Water quality parameters monitored throughout test and remained withing acceptable limits. Samples analyzed for concentration of test article varied widely. This variability was attributed to the instability of the test compound in water and to incomplete dispersion. Nominal concentrations of test compound were 0.00, 0.03, 0.06, 0.12, 0.25 and 0.50 mg/l. LC50s were recorded at 24, 96 and 336 hours. At the time the test was terminated, no mortalities had occurred during the preceeding 48 hours.

Reference: Monsanto SR-80-1803058-A1, SRI International, 1981

# 5. TOXICITY

#### \*5.1 ACUTE TOXICITY

#### 5.1.1 ACUTE ORAL TOXICITY

Type:  $LD_0[]; LD_{100}[]; LD_{50}[X]; LDL_0[]; Other[]$ 

Species/strain: Sprague-Dawley Albino Rats

Value: 730 mg/kg b.w.

Discriminating dose: 794 mg/kg

Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [x] Klimisch 2
Test substance: Santoflex 77, Lot # KC01-04, purity:>94%

Remarks: Groups of male and female rats were fed either 501, 631, 704 or 1000

mg/kg of the undiluted test substance as a single oral dose by gavage. Clinical signs of toxicity included reduced appetite and activity – for to six days in survivors – followed by increasing weakness, collapse and death. Gross autopsy findings on decedents included hemorrhagic areas of the lungs, liver discoloration and acute gastrointestinal inflammation. Survivors were sacrificed after 10 days. All viscera examined appeared

normal.

Reference: Monsanto Y-73-168 Younger Laboratories, 1973

# 5.1.2 ACUTE INHALATION TOXICITY

Type:  $LC_0[X]; LC_{100}[]; LC_{50}[]; LCL_0[]; Other[]$ 

Species/strain: Sprague-Dawley Albino rats

Exposure time: 6 hours w/10 day observation period

Value: Sample did not vaporize

Method: Ambient Temperature Inhalation

GLP: Yes [ ] No [ ] ? [x] Klimisch 2
Test substance: Santoflex 77 Lot # KC01-04, purity: >94%

Remarks: Male rats were exposed to the test article in an inhalation chamber for a

period of six hours at ambient temperature. The initial sample size of the test article was 133 grams. At the end of six hours, the sample was reweighed and found to be 133 grams, and no sample was recovered from the chamber air condenser. Santoflex 77 did not vaporize under the test conditions. No animal experienced any symptoms of toxicity. The 10 day observation period was uneventful, and all animals survived to sacrifice with no noted ill-effects. Autopsy findings were that all viscera examined

appeared normal.

Reference: Monsanto Y-73-168 Younger Laboratories, 1973

#### 5.1.3 ACUTE DERMAL TOXICITY

Remarks:

Type:  $LD_0[\ ]; \ LD_{100}[\ ]; \ LD_{50}[\ X]; \ LDL_0[\ ]; \ Other[\ ]$ 

Species/strain: New Zealand Albino Rabbits

Value: >3160 mg/kg b.w. Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex 77, Lot # KC01-04, purity: >94%

The undiluted test substance was applied to the shaved skin of male and female rabbits for a period of 24 hours, followed by a 14 day recovery period. Dosages were 1260, 2000, 3160, 5010 or 7940 mg/kg. Clinical signs of toxicity were reduced appetite and activity – three to seven days in survivors – followed by increasing weakness, collapse and death. Gross autopsy findings on decedents included lung hyperemia, liver discoloration, enlarged gall bladder and gastrointestinal inflammation. Survivors were sacrificed following the recovery period. All viscera appeared normal on all but two animals, which exhibited a slight discoloration of both liver and

kidneys.

Reference: Monsanto Y-73-168 Younger Laboratories, 1973

#### 5.2 CORROSIVENESS/IRRITATION

# 5.2.1 SKIN IRRITATION/CORROSION

Species/strain: New Zealand Albino Rabbits

Results: Highly corrosive [ ]; Corrosive [ ]; Highly irritating [ ];

Irritating [ ]; Moderate irritating [ ]; Slightly irritating [ ];

Not irritating [X]

Classification: Highly corrosive (causes severe burns) [ ];

Corrosive (causes burns) [ ]; Irritating [ ]; Not irritating [X]

Method: Primary Skin Irritation

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex 77 Lot #KC01-04, purity:>94%

Remarks: 0.5 ml of the undiluted test substance was applied to the shaved skin of six

male and female rabbits. Irritation was scored on a scale of 0-4 for both erythema and edema. The 24 hour score for all animals was 0.0, indicating the test substance was non-irritating. Observations noted was a slight defatting effect on the skin, with mild flaking after 7-10 days.

Reference: Monsanto Y-73-168 Younger Laboratories, 1973

# 5.2.2 EYE IRRITATION/CORROSION

Species/strain: New Zealand Albino Rabbits

Results Highly corrosive [ ]; Corrosive [ ]; Highly irritating [ ];

Irritating [ ]; Moderate irritating [ ]; Slightly irritating [X];

Not irritating [ ]

Classification: Irritating [X]; Not irritating []; Risk of serious damage to eyes []

Method: Draize

GLP: Yes [] No [] ? [X] Klimisch 2
Test substance: Santoflex 77 Lot # KC01-04, purity: >94%

Remarks: 0.1 ml of the undiluted test substance was applied to the eyes of rabbits.

Irritation was assessed at 1, 24, 48, 72 and 168-hour intervals on the basis of irritation to the cornea, iris and conjunctivae. Immediate findings were slight discomfort. 1-hour findings were slight erythema, very slight edema and copious discharge. 24-hour score was 10.0, 48-hour score was 9.3, 72-hour score was 6.3 and 168-hour score was 0.0. The 24/48/72 hour average score was 8.5 for a classification as a "slight" acute eye irritant.

Reference: Monsanto Y-73-168 Younger Laboratories, 1973

# 5.3 SKIN SENSITISATION

Type:

Species/strain:

Results: Sensitizing [ ]; Not sensitizing [ ]; Ambiguous [ ] Classification: (if possible, according to EC Directive 67/548/EEC)

Sensitizing [ ]; Not sensitizing [ ]

Method:

GLP: Yes[] No[]?[]

Test substance: ...., purity:....

Remarks: Reference:

#### \*5.4 REPEATED DOSE TOXICITY

Species/strain: Sprague-Dawley CD Rats

Sex: Female [ ]; Male/Female [ X ]; No data [ ]

Route of Administration: Oral/Dietary

Exposure period: 30 days
Frequency of treatment: Daily
Post exposure observation period:

Dose: 0, 100, 300, 500, 1000 and 2000 ppm

Control group: Yes [ X ]; No [ ]; No data [ ];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL: 100 ppm for males, 300 ppm for females

LOEL: Not Determined

Results: In a 30-day range-finding study that preceded a 90-day study, the test

substance was administered orally, via dietary admixture, to groups of male and female rats (5/sex/group). Control animals received the standard laboratory diet. Physical observations, body weight and food consumption measurements were performed on all animals pretest and at selected intervals during the study. Hematology and chemistry determinations were performed on all animals at study termination. There were no mortalities during the course of the study. After four weeks of treatment, all animals were sacrificed, selected organs were weighed, and organ/body weight ratios were calculated. Complete postmortem examinations were conducted on all animals. Differences from control in mean body weights were statistically significant at 500 ppm and 1000 ppm males and in 2000

ppm males and females. Differences from control in mean body weight/body weight gain suggested a treatment-related effect in males at dose levels at and above 300 ppm, and in females at and above 1000 ppm. Food consumption values in Week 1 were reduced for males at 500 ppm and above, and for females at 300 ppm and above. Food consumption at Weeks 3-4 was comparable to controls. Males and females at the two highest dose levels exhibited increased mean platelet counts following four weeks of treatment. Males in these groups also exhibited increased mean erythrocyte. The mean hematology values for males and females in all treatment groups were comparable to controls. Alterations in several clinical chemistry parameters were noted for higher dose levels. Mean terminal body weights were reduced at the two highest dose levels in females, and at the three highest dose levels in males. While several organs in treated males and females exhibited alterations in either mean absolute or relative weights, these changes were considered secondary effects and not indicative of significant organ toxicity. Gross pathological examination did not reveal any effects that were considered treatment-

related.

Method: Dunnett, C.W., A Multiple Comparison Procedure for

Comparing Several Treatments with a Control, Jour. Am. Stat.

Assoc. 50: 1096-1121, 1955

Yes [X] No [] ? [] GLP: Klimisch 1 Santoflex 77 Lot# KJ01-03, purity: 99+% active Test substance:. Reference: Monsanto BD-87-146 Bio/dynamics Labs, 1987

Species/strain: Sprague-Dawley CD Rats

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral/Dietary

Exposure period: 90 days Daily Frequency of treatment: Post exposure observation period:

Dose: Males: 0, 100, 250 and 500 ppm Females: 0, 250, 500 and 750 ppm

Control group: Yes [ **X** ]; No [ ]; No data [ ];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL: 100 ppm for males, not established for females

LOEL: Not Determined

The test substance was administered orally, via dietary admixture, to Results:

> groups of male and female rats (10/sex/group). Control animals received the standard laboratory diet. Physical observations, body weight and food consumption measurements were performed on all animals pretest and at selected intervals during the study. Hematology and chemistry determinations were performed on all animals at Months 1.5 and 3. There were no mortalities during the course of the study. After three months of treatment, all animals were sacrificed, selected organs were weighed, and organ/body and organ/brain weight ratios were calculated. Complete were postmortem examinations conducted on all animals. Histopathological evaluation of selected tissues was performed on all control and high-dose animals. The lungs, spleen, liver and kidneys were examined microscopically for all animals in all groups. Mean body weights and mean body weight gains were reduced in males at 250 and 500 ppm,

and in all treated females. Overall, mean food consumption values for all treated groups were comparable to controls. Several clinical chemistry parameters exhibited statistically significant differences from control. Alkaline phosphatase was elevated in the 500 ppm males and 750 ppm females at Month 3. Mean serum glutamic oxaloacetic transaminase levels were significantly reduced in the 100, 250 and 500 ppm males at Month 1.5 but not at Month 3. Mean serum glutamic pyruvic transaminase was reduced in the 500 and 750 ppm females at Month 3. Several organs in the treated males and females exhibited alterations in mean absolute and/or relative (to body or brain) weight data. However, these alterations were generally consistent with the reductions noted in body weight data and were considered secondary effects which were not considered indicative of significant organ toxicity. There were no treatment-related findings noted in mortality, physical observations, opthalmoscopic, hematology, organ weight or gross and microscopic pathology.

OECD Guidelines for Testing of Chemicals, Section 453, 1981 and Method:

USEPA TSCA Section 4(a) Test Rules, 1982

GLP: Yes [X] No [] ? [] Klimisch 1 Santoflex 77 Lot# KJ01-03, purity: 99+% active Test substance:. Reference: Monsanto BD-87-147 Bio/dynamics Labs, 1989

Species/strain: Charles River Albino rats

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral/Dietary

Exposure period: 2 years Frequency of treatment: Daily Post exposure observation period:

Dose: 0, 30, 100 or 300 ppm

Control group: Yes [X]; No []; No data [];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL: 30 ppm LOEL: 100 ppm

A two-year chronic oral toxicity study was conducted on groups of 400 Results:

CD Outbred rats (50/sex/dose) at dietary levels ranging from 0-300 ppm. Reductions in body weights and body weight gains were noted for males and females at the 300 ppm dose throughout the investigation. Body weights of females fed 100 ppm were reduced during the first 7 weeks, and for 100 ppm males for the first 4 weeks. After those intervals, body weights compared favorably with controls. 30 ppm animals had body weights and weight gains that compared favorably with controls. Frequency and distribution of deaths during the investigation for all dose levels was similar to controls. Gross pathological examination of animals that died during the study did not reveal any relation between death and exposure to the test substance. No unusual behavioral reactions were noted in dosed animals during the course of the study. Results of hematologic studies conducted - total and differential leukocyte count, erythrocyte count, hemoglobin concentration, hematocrit value, mean corpuscular volume, mean corpuscular hemoglobin and mean corpuscular hemoglobin concentration – were either similar to, or within the range of expected values for this strain of albino rats of this age and in this laboratory. Results of clinical blood chemistry studies (SGPT, BUN, SGOT, Fasting Blood Glucose Concentration, SAP) and of urinalyses (glucose, albumin, microscopic elements, pH and specific gravity) conducted showed similar results between control and test animals. Gross pathological examinations of animals sacrificed at 24 months revealed similar findings between test and control animals. Histopathological examinations of tissues and organs from the control and 300 ppm animals sacrificed at 24 months showed no treatment-related lesions. Microscopic examination of suspect neoplasms among all sacrificed animals and all animals that died during the study were conducted. No differences were noted between test and control rats as to the organ system involved, the type or the classification of neoplasms. The spectrum of neoplasms observed compared favorably to historical data at this laboratory for rats of this strain and age. At 17.5 months of testing, tetracycline HCl was added to the diets of all groups (30g/kg of diet) for a two-week period to treat a severe respiratory infection which caused an increase in mortality in both control and treated animals.

Method: 2-Year Chronic Oral Toxicity IBT Protocol # 622-05400B (1974)

GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex 77 reddish liquid Lot# KD05-57, purity: 99+% active

Reference: Monsanto BTL-74-27, Industrial Bio-Test Labs, 1978

#### \*5.5 GENETIC TOXICITY IN VITRO

#### A. BACTERIAL TEST

Type: Ames Reverse Bacterial Mutation

System of testing: Salmonella typhimurium TA98, TA100, TA1535, TA1537 Concentration: 0.01, 0.04, 0.2, 1, 3, 10, 40 and 200 micrograms/plate

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 200 micrograms/plate

Without metabolic activation: 10 micrograms/plate

Precipitation conc: 1 microgram/plate

Genotoxic effects: + ?

With metabolic activation: [][][X]
Without metabolic activation: [][][X]

Method: Ames, B.N., McCann, J. and Yamaski, E. Methods for Detecting

Carcinogens and Mutagens with the Salmonella Mammalian-Microsome

Test. Mutat. Res. 31, 347-364, 1975

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 77 Lot# 7/31/85WGK, purity: 99+% active

Remarks: Santoflex 77 was tested in Ames/Salmonella plate incorporation assays

using the tester strains TA98, TA100, TA1535 and TA1537 in the presence and absence of an Aroclor-induced rat liver mammalian metabolic activation system (S-9 Mix). No mutagenic activity was observed for the test compound in any of these assays. Toxicity of the test

compound was significantly reduced in the presence of the S-9 Mix.

Reference: Monsanto ML-85-242 Environmental Health, 1985

# B. NON-BACTERIAL IN VITRO TEST

Type: Mammalian Cell Gene Forward Mutation Assay

System of testing: L5178Y Mouse Lymphoma cells

Concentration: 0.002, 0.004, 0.008, 0.016 (without activation) 0.002, 0.004, 0.008, 0.016, 0.032 (with activation)

0.002, 0.004, 0.008, 0.016, 0.032 (with activation)

Metabolic activation: With [ ]; Without [ ]; With and Without [ X ]; No data [ ]

Results:

Cytotoxicity conc: With metabolic activation: 0.032 ug/ml

Without metabolic activation: 0.016 ug/ml

Precipitation conc: Not determined

Genotoxic effects: + ? -

With metabolic activation: [][][X]
Without metabolic activation: [][][X]

Method: Clive and Spector, Mutation Research 31:17-29 (1975)

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: CP-25477 (Santoflex 77) dark liquid, purity >94%

Remarks: The test article was evaluated for specific locus forward mutation in the

L5178Y Thymidine Kinase (TK) mouse lymphoma cell assay. Stock solutions were prepared in DMSO. DMSO was used as the negative control. EMS was used as the positive control without activation and DMN was used as the positive control with activation. The test article was

found to be negative

Reference: Monsanto BIO-76-246 Litton Bionetics, 1976

Type: <u>In vitro</u> Unscheduled DNA Synthesis (UDS)

System of testing: Primary rat hepatocyte cultures (Fischer-344 strain)

Concentration: 0.01, 0.05, 0.1, 0.5, 1, 5, 10, 20, 50, 100, 500, 1000 ug/ml Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: Preliminary Assay: 50 ug/ml

Replicate Assay: 5 ug/ml

Precipitation conc: Separation (two layers) at 1000 ug/ml

Genotoxic effects: + ? -

[][][X]

Method: Williams, G.M., Detection of Chemical Carcinogens by Unscheduled

DNA Synthesis in Rat Liver Primary Cell Cultures, Cancer Research 37,

pp. 1845-1851 (1977)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 77 liquid produced 07/31/85, purity 99+% active

Remarks: Acetone (1%) used as solvent and diluent. Primary rat liver cell cultures

derived from the livers of two adult male rats. The positive control was 2-AAF, the solvent control was acetone in the preliminary assay and DMSO in the replicate assay. The percentage of cells in repair was calculated as the percentage of cells with at least 5 net grains/nucleus. 150 cells were scored for each concentration reported for each experiment. The net grain counts were negative at each concentration of the test compound, in the solvent control and in the medium control, in contrast to the strong positive response produced by the positive control 2-AAF in both experiments. These results indicate that Santoflex 77 is not a genotoxic agent under the

conditions of the in vitro rat hepatocyte DNA repair assay.

Reference: Monsanto SR-85-250, SRI International, 1986

# \* 5.6 GENETIC TOXICITY IN VIVO

Type:

Species/strain:

Sex: Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]

Route of Administration:

Exposure period:

Doses: Results:

Effect on mitotic index or P/N ratio:

Genotoxic effects: + ? -

[][][]

Method:

GLP: Yes [ ] No [ ] ? [ ]

Test substance: Remarks: Reference:

#### 5.7 CARCINOGENICITY

Species/strain:

Sex: Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]

Route of Administration:

Exposure period:

Frequency of treatment:

Postexposure observation period:

Doses:

Control group: Yes [ ]; No [ ]; No data [ ];

Concurrent no treatment [ ]; Concurrent vehicle [ ]; Historical [ ]

Results: Method:

GLP: Yes [] No [] ? []

Test substance:` ...., purity:....

Remarks: Reference:

#### \*5.8 TOXICITY TO REPRODUCTION

Type: Fertility [X]; One-generation study []; Two-generation study [];

Other [X] Three Generation Study

Species/strain: Charles River Albino Rats

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral/Dietary

Exposure period: Premating, throughout mating, gestation and lactation

Frequency of treatment: Daily

Post exposure observation period: Not Determined

Premating exposure period: male: F0 - 14 wks F1 - 14 wks F2 - 18 wks

female: F0 - 14 wks F1 - 14 wks F2 - 18 wks

Duration of the test: F0 - 23 wks F1 - 23 wks F2 - 26 wks

Doses: 0, 30, 100 or 300 ppm

Control group: Yes [X]; No []; No data [];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL Parental: 30 ppm (based on reduced body weight gain)

NOEL F1 Offspring: 30 ppm (based on reduced pup survival) NOEL F2 Offspring: 30 ppm (based on reduced pup survival)

Results: Santoflex 77 was administered to three successive generations of rats at

dose levels of 0, 30, 100 or 300 ppm. Dose levels were selected on the basis of results from a previous 2-year chronic oral feeding study. No adverse effects on mating or fertility indices were noted in any of the treated animals. Reduced survival of offspring was observed in the mid-to high-dose groups. Evidence of parental toxicity was also present as

indicated by reduced body weights of mid-to high-dose animals

General parental toxicity: Reduced body weights and mean body weight gains were noted for the 100 and 300 ppm males and females. No other treatment-related

effects were evident in results of clinical blood chemistry studies and urinalyses between the control groups and the treated animals.

Toxicity to offspring: A small but statistically significant reduction in the survival rates of pups was noted in the 100 ppm and 300 ppm groups.

Method: 3-Generation Reproductive Toxicity IBT Protocol # 622-05400C (1974)

GLP: Yes [ ] No [ ] ? [ **X** ] Klimisch 2

Test substance: Santoflex 77 dark red liquid Lot# KD05-57, purity: 99+% active

Remarks: Protocol similar to Monsanto BTL-74-27, Industrial Bio-Test Labs, 1978

Reference: Monsanto BTL-76-145, Industrial Bio-Test Labs, 1976

# \*5.9 DEVELOPMENTAL TOXICITY/ TERATOGENICITY

Species/strain: Charles River CD Albino Rats

Sex: Female [ X ]; Male [ ]; Male/Female [ ]; No data [ ]

Route of Administration: Oral gavage

Duration of the test: 25 days from mating to last C-section

Exposure period: Day 6-15 of gestation

Frequency of treatment: Daily, as a single oral dose at a volume of 5 ml/kg

Doses: 25, 75 and 150 mg/kg/day Control group: Yes [ X ]; No [ ]; No data [ ];

Concurrent no treatment [ ]; Concurrent vehicle [X]; Historical [ ]

NOEL Maternal Toxicity: 25 mg/kg/day NOEL teratogenicity: 150 mg/kg/day

Results: Groups of 25 mated CD rats were assigned to one control group and three

treatment groups to determine the teratogenic potential of the test substance. Dosage levels of 25, 75 and 150 mg/kg/day were administered orally by gavage as a single daily dose on Days 6-15 of gestation. The control group received the corn oil vehicle only. Cesarean sections were performed on all surviving females on gestation Day 20, and the fetuses

removed for teratologic evaluation.

Maternal general toxicity: Toxicity in the dams was apparent at the 75 and 150 mg/kg/day dosage levels. Parameters adversely affected were maternal survival, appearance, behavior and body weight gain. Four of the 150 mg/kg/day females and one 75 mg/kg/day female died between gestation Days 16-17. Control animals and the low dose group had 100% survival. Antemortem abnormalities in the decedents included dried blood around and/or expelled from the vaginal orifice, blood under the cage, stained, wet or matted coat, hypothermia and ptyalism. There were no treatment-related gross internal lesions evident. No effect on Cesarean section observations

was noted in the dams at any dosage level.

Pregnancy/litter data: No obvious differences were noted between the

Treated groups and the control group.

Foetal data: Malformations that were observed in the treated groups occurred in low incidence and were not considered treatment-related. One high-dose fetus had anophthalmia, one mid-dose and two control group fetuses had microphthalmia, and another mid-dose fetus had ectopia cordia and sternoschisis. There were no adverse effects on the fetal parameters examined (survival, growth, morphological development) at dose levels at or below 150 mg/kg/day.

OECD Guidelines for Testing of Chemicals No. 414 "Teratogenicity"

1981, and TSCA Health Effects Guidelines "Teratogenicity Study" 1982

GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex 77 red-brown liquid Lot# 25477, purity: 99+% active

Remarks: Based on the results, the test article did not induce developmental toxicity

In the offspring of Charles Rived CD rats under the test conditions.

Reference: Monsanto IR-85-290 International Research and Development, 1986

#### 5.10 OTHER RELEVANT INFORMATION

# A. Specific toxicities

Method:

Type: Immunotoxicity – Repeated Insult Patch Testing

Modified Schwartz Method and Shelanski Method

Results: Several studies were run using human volunteers to determine the potential

for Santoflex 77 to cause allergic skin reactions in compounded rubber stocks. Loading of the test article was from 0.5 to 3 phr (parts per hundred rubber) in a typical B-1 Masterbatch. Some study results indicated that the test article caused no primary irritation and no allergic response, while other study

results were positive for sensitization.

Remarks: Differences in responses may be due to the presence of other chemicals in

the B-1 masterbatch formulations.

Reference: Monsanto SH-61-17, Industrial Biology Labs, 1961

Monsanto SH-63-10, Industrial Biology Labs, 1963 Monsanto SH-64-4, Industrial Biology Labs, 1964 Monsanto SH-64-5, Industrial Biology Labs, 1964 Monsanto SH-73-12, Industrial Biology Labs, 1973

#### B. Toxicodynamics, toxicokinetics

Type: (e.g. toxicodynamics, toxicokinetics)

Results: Remarks: References:

#### \* 5.11 EXPERIENCE WITH HUMAN EXPOSURE

Results: Remarks: Reference:

#### 6. REFERENCES

- 1. United States National Toxicology Program, November 6, 1990
- 2. Monsanto Physical Constants of CP25447. Standard Manufacturing Process Manual, July 1977
- 3. American Society for Testing and Materials, 1997
- 4. Monsanto SRI 8669, Selected Environmental Fate Studies of Nine Chemical Compounds, SRI International, August 20, 1980

- 5. USEPA federal Register Volume 44, No. 53, March 16, 1979, pp. 16 and 255
- 6. American Society for Testing and Materials, D 92, Standard Test Method for Flash and Fire Points by Cleveland Open Cup, 1997
- 7. Monsanto Report ABC-32303, Santoflex 77 Phase I Hydrolysis Study: Identification of Hydrolysis Products, Analytical BioChemistry Laboratories, January 15, 1986
- 8. Monsanto ES-79-SS-25, Environmental Persistence Screening of Selected Rubber Chemicals, Monsanto Industrial Chemicals Environmental Sciences, December 28, 1979
- 9. American Society for Testing and Materials, Draft Method No. 2, ASTM Committee E35.24, August 1979
- 10. Monsanto BN-76-254 Acute (96 Hour) Toxicity of Santoflex 77 to Rainbow Trout, EG&G Bionomics Aquatic Toxicity Laboratory, December 1976
- 11. Monsanto BN-76-254 Acute (96 Hour) Toxicity of Santoflex 77 to Bluegill Sunfish, EG&G Bionomics Aquatic Toxicity laboratory, December 1976
- 12. Monsanto AB-79-1384361-1b Acute Toxicity of Santoflex 77 to <u>Daphnia magna</u>, Analytical BioChemistry Laboratories, August 27, 1979
- 13. Monsanto AB-79-1384361-1a Acute Toxicity of Santoflex 77 to Fathead Minnows, Analytical BioChemistry Laboratories, August 27, 1979
- 14. Monsanto BN-79-1384361-2 Toxicity of Santoflex 77 to the freshwater alga <u>Selenastrum</u> capricornutum, EG&G Bionomics Marine Research Laboratory, August 1979
- 15. Monsanto AB-81-9AB981014, Acute Toxicity of Santoflex 77 to Midge, Analytical BioChemistry Laboratories, August 19, 1981
- Gettings, A.V and W.J. Adams. 1980. Method for Conducting Acute Toxicity Tests with the Midge <u>Paratanytarsus parthenogenetica</u>. Monsanto Industrial Chemicals Company, Report ES-81-M-1
- 17. C.E. Stephan, Chairman, Committee on Methods for Toxicity Tests with Aquatic Organisms, US EPA, 1975
- 18. Monsanto AB-80-1803058-B1, Flow-Through Bioassay Final Report: Dynamic Acute Toxicity of Santoflex 77 to Fathead Minnows, Analytical BioChemsitry Laboratories, January 20, 1981
- 19. Monsanto SR-80-1803085-A1, Time Independent Toxicity Study on Santoflex 77 using Fathead Minnows as the Test Organism, SRI International, September 8, 1981
- 20. Monsanto Y-73-168, Toxicological Examination of CP-25477 (Santoflex 77) for Acute Oral and Dermal Toxicity, Younger Laboratories, October 9, 1973
- 21. Monsanto Y-73-168, Toxicological Examination of CP-25477 (Santoflex 77) for Ambient Temperature Inhalation Toxicity, Younger Laboratories, October 9, 1973
- 22. Monsanto Y-73-168, Toxicological Examination of CP-25477 (Santoflex 77) for Acute Eye and Primary Skin Irritation, Younger Laboratories, October 9, 1973
- 23. Monsanto BD-87-146, A 4 Week Range-Finding Toxicity Study with Santoflex 77 in the Rat Via Dietary Admixture, Bio/dynamics, Inc. June 14, 1989
- 24. Monsanto BD-87-147, A Subchronic 3-Month Oral Toxicity Study with Santoflex 77 in the Rat Via Dietary Admixture, Bio/dynamics, Inc. April 28, 1989
- 25. Monsanto BTL-74-27, Two-Year Chronic Oral Toxicity Study with Santoflex 77 in Albino Rats, Industrial Bio-Test Laboratories, Inc. November 27, 1978
- 26. Monsanto ML-85-242, Ames/<u>Salmonella</u> Mutagenicity Assay of Santoflex 77, Monsanto Environmental Health Laboratory, February 18, 1986
- 27. Monsanto SR-85-250, Evaluation of the Potential of Santoflex 77 to Induce Unscheduled DNA Synthesis in Primary Rat Hepatocyte Cultures, SRI International, May 23, 1986
- 28. OECD Guidelines for Testing of Chemicals; No. 414, Teratogenicity, adopted May 1981
- US EPA Report 560/6-82-001, TSCA Health Effects Test Guidelines for Teratogenicity Studies, August 1982
- 30. Monsanto IR-85-290, Teratology Study in Rats with Santoflex 77, International Research and Development Corporation, April 1, 1986

- 31. Monsanto SH-61-17, Repeated Insult Patch Tests of Antidegradants, Industrial Biology Laboratories, Inc. May, 1961
- 32. Monsanto SH-63-10, Modified Schwartz Patch Test Study of Monsanto Rubber Samples, Industrial Biology Laboratories, Inc., November 8, 1963
- 33. Monsanto SH-64-4, Repeat Insult Patch Test on Vulcanized Rubbers, Industrial Biology Laboratories, May 5, 1964
- 34. Monsanto SH-64-5, Dermatitic Studies of Hexyl- and Heptyl-PPDs in Rubber, Industrial Biology Laboratories, March 1964
- 35. Monsanto SH-73-12, Repeat Insult Patch Test with Uncured Rubbers, Industrial Biology Laboratories, April 1973

# 3081-01-4

# p-Phenylenediamine, N-(1,4-dimethylpentyl)-N'-phenyl-

# 2. PHYSICAL-CHEMICAL DATA

#### \*2.1 MELTING POINT

Value: 32.4°C for highly purified (99+%)

Otherwise, room temperature viscous liquid

Decomposition: Yes [ ] No [X] Ambiguous [ ] Sublimation: Yes [ ] No [X] Ambiguous [ ]

Method: Crystallizing Point

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Remarks: Physical Constants, Flexsys SMP, R.L. Wright (1982)
Reference: Flexsys 7PPD Standard Manufacturing Process

\*2.2 BOILING POINT

Value: 231 °C

Pressure: at 3.5 mm Hg

Decomposition: Yes [] No [X] Ambiguous []

Method: Instrumental – Differential Scanning Calorimeter (DSC)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Remarks: Physical Constants, Flexsys SMP, R.L. Wright (1982)

Reference: L.M. Baclawski Notebook #2355311 (1982)

†2.3 DENSITY (relative density)

Type: Bulk density []; Density [X]; Relative Density []

Value: 1.0 Temperature: 20 °C

Method: Flexsys Standard Method of Analysis FF97.4-1

GLP: Yes [X] No[] ?[]

Remarks: Hydrometer method. Hydrometer must meet standards set in

ASTM-E-100

Reference: Flexsys 7PPD Standard Manufacturing Specifications

\*2.4 VAPOUR PRESSURE

Value: 1.25 x 10(-10) mm Hg

Temperature: 25 °C

Method: calculated [X]; measured [ ]

Antoine Equasion.

GLP: Yes [ ] No [X] ? [ ] Klimisch 2

Remarks: None

Reference: Monsanto Toxicology Profile, Santoflex 14, C.E. Healy 1993

\*2.5 PARTITION COEFFICIENT log<sub>10</sub>P<sub>ow</sub>

Log Pow: 5.17

Temperature: Not Applicable

Method: calculated [X]; measured [ ] Klimisch 2

SRC LogKow (KowWin) Program 1995

GLP: Yes [] No [X] ? []

Remarks: None

Reference: Meylan, W.M. and. P.H. Howard, 1995 J. Pharm. Sci. 84: 83-92

# \*2.6 WATER SOLUBILITY

A. Solubility

Value: 0.67 mg/l in pH 7.0 deionized water

Temperature: 25°C

Description: Miscible []; Of very high solubility [];

Of high solubility []; Soluble []; Slightly soluble [];

Of low solubility []; Of very low solubility [X]; Not soluble []

Method: Saturated Solution/GC Analysis

GLP: Yes [X] No [] ? [] Klimisch 1

Remarks: Preliminary solubility study for Phase I Hydrolysis

Reference: Monsanto ABC 32305, Analytical Bio-Chemistry Labs, 1986

B. pH Value, pKa Value

pH Value: Not Applicable

Concentration: Temperature: Method: .

GLP: Yes [] No [] ? []

pKa value Remarks: Reference:

2.11	OXIDISING PROPERTIES			
	Results:	Maximum burning rate equal or higher than reference mixture[];		
		Vigorous reaction in preliminary test [ ];		
		No oxidising properties [X]; Other [ ]		
	Method:			
	GLP:	Yes [ ] No [ ] ? [ ]		
	Remarks:			
	Reference:			
†2.12	OXIDATION: F	REDUCTION POTENTIAL		
	Value:	Not Applicable		
	Method:			
	GLP:	Yes [ ] No [ ] ? [ ]		
	Remarks:			
	Reference:			
2.13	ADDITIONAL 1	DATA		
A.	Partition co-efficient between soil/sediment and water (Kd)			
	Value:			
	Method:			
	GLP:	Yes [ ] No [ ] ? [ ]		
	Remarks:			
	Reference:			
В.	Other data			
	Results:			
	Remarks:			
	Reference:			

## 3. ENVIRONMENTAL FATE AND PATHWAYS

## \*3.1.1 PHOTODEGRADATION

Type: Air [X]; Water [ ]; Soil [ ]; Other [ ] Light source: Sunlight [ ]; Xenon lamp [ ]; Other [ ]

Light spectrum: .....nm

Relative intensity: ..... (based on intensity of sunlight)

Direct photolysis:

Half life: .....

Degradation: ...... % (weight/weight) after ..... (exposure time)

Quantum yield: .....

Indirect Photolysis:

Type of sensitizer: ..... OH ...

Degradation: ..... 50% at 0.563 Hrs...

Method: calculated [X]; AOP Program (v1.89)

measured [ ]

GLP: Yes [ ] No [ X ] ? [ ]

Test substance: molecular structure, purity:.....

Remarks:

Reliability: (2) valid with restrictions

Accepted calculation method

Reference: Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

## \*3.1.2 STABILITY IN WATER

Type: Abiotic (hydrolysis) [X]; biotic (sediment)[ ]

Half life: Not Measured

Degradation: 96% at pH 7.0 at 25 °C after 24 Hours
Method: Extraction, ABC Protocol M-8305 (1986)
GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex 14 purple liquid, Lot# KD09-813, purity:>95%

Remarks: No test substance detected at seven days. Hydrolysis products

identified by GC analysis as 4-hydroxydiphenylamine (35%) and Benzoquinoneimine-n-phenyl (65%). Stock solution in acetone.

Reference: Monsanto ABC 32305, Analytical Bio-Chemistry Labs, 1986

#### \*3.2 MONITORING DATA (ENVIRONMENTAL) ..... Type of Measurement: Background [ ]; At contaminated site [ ]; Other [ ] Media: Results: Remarks: Reference: 3.3 **TRANSPORT AND DISTRIBUTION** BETWEEN **ENVIRONMENTAL** COMPARTMENTS **INCLUDING ESTIMATED ENVIRONMENTAL** CONCENTRATIONS AND DISTRIBUTION \*3.3.1 **TRANSPORT** Adsorption [ ]; Desorption [ ]; Volatility [ ]; Other [ ] Type: Media: Method: Results: Remarks: Reference: \*3.3.2 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION) Media: Air-biota []; Air-biota-sediment-soil-water []; Soil-biota []; Water-air []; Water-biota []; Water-soil []; Other [] Method: Fugacity level I [ ]; Fugacity level II [ ]; Fugacity level III [X]; Fugacity level IV [ ]; Other (calculation) [ ] Results: С Half-Life Concentration Emissions Fugacity (percent) (hr) (kg/hr) (atm) Air 0.027 1.13 7.19e-013 1000 Water 15.2 900 1000 3.5e-014 Soil 57.5 900 1000 1.11e-015 Sediment 27.2 3.6e + 0032.36e-014 Reaction Advection Reaction Advection (kg/hr) (kg/hr) (percent) (percent) 531 8.64 17.7 0.288 Air Water 375 487 12.5 16.2 47.1 Soil 1.41e+003 0 Λ 17.4 0.58 Sediment 168 5.58 Persistence Time: 1.06e+003 hr Reaction Time: 1.28e+003 hr Advection Time: 6.23e+003 hr Percent Reacted: Percent Advected: 17.1 Remarks: Reliability: (2) valid with restrictions Accepted calculation method

Reference:

Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

## \*3.5 BIODEGRADATION

Type: aerobic [X]; anaerobic [ ]
Inoculum: adapted [X]; non-adapted [];

Concentration of the chemical: 20.0 mg/l related to COD [X]; DOC[]; test substance[] Medium: water []; water-sediment []; soil []; sewage treatment [X]

Degradation: 0 % after 35 days

Results: readily biodeg. []; inherently biodeg. []; under test condition no

biodegradation observed [X], other [ ]

Kinetic

Method: ASTM Draft 3 Proposed Standard Practice for the Determination

Of the Ultimate Biodegradation of Organic Chemicals (1980).

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 14 purple liquid Lot#KA01-07, purity: >95%

Remarks: Shake Flask carbon dioxide evolution test. Glucose and Sodium

Citrate used as positive controls.

Reference: Monsanto ES-80-SS-48 MIC Environmental Sciences 1981

## 4. <u>ECOTOXICITY</u>

## \*4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [ X ]

Species: Salmo gairdneri (Rainbow Trout)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = >1.00 mg/l

 $LC_{50}$  (48h) = 0.70 mg/l  $LC_{50}$  (72h) = Not Determined  $LC_{50}$  (96h) = 0.42 mg/l NOEC = 0.18 mg/l LOEC = Not Determined

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 14 purple liquid, purity:>95%

Remarks: Stock solutions prepared in nanograde Acetone. Water quality

parameters (pH, temperature, dissolved oxygen content) monitored throughout test. Test fish challenged with Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto ABC 30687, Analytical Bio-Chemistry Labs, 1983

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [  ${f X}$  ]

Species: Lepomis macrochirus (Bluegill Sunfish)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 0.38 mg/l

 $LC_{50}$  (48h) = 0.30 mg/l  $LC_{50}$  (72h) = Not Determined  $LC_{50}$  (96h) = 0.30 mg/l NOEC = 0.18 mg/l

LOEC = 0.18 mg/r

LOEC = Not Determined

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 14 purple liquid, purity:>95%

Remarks: Stock solutions prepared in nanograde Acetone. Water quality

parameters (pH, temperature, dissolved oxygen content) monitored throughout test. Test fish challenged with Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto ABC 30686, Analytical Bio-Chemistry Labs, 1983

Type of test: static [X]; semi-static []; flow-through []; other []

open-system [ ]; closed-system [ X ]

Species: <u>Pimephales promelas</u> (Fathead Minnows)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 1.30 mg/l

 $LC_{50}$  (48h) = 1.30 mg/l  $LC_{50}$  (72h) = Not Determined  $LC_{50}$  (96h) = 1.10 mg/l NOEC = 0.32 mg/l LOEC = Not Determined

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 14 purple liquid, purity: >96%

Remarks: Stock solutions prepared in nanograde Acetone. Water quality

parameters (pH, temperature, dissolved oxygen content) monitored throughout test. Test fish challenged with Antimycin

A. Data reported at 95% confidence level.

Reference: Monsanto ABC 31116, Analytical Bio-Chemistry Labs, 1983

## 4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

## \*A. Daphnia

Type of test: static [X]; semi-static []; flow-through []; other [];

open-system [ ]; closed-system [ X ]

Species: <u>Daphnia magna</u>

Exposure period: 48 Hours

Results:  $EC_{50}$  (24h) = 0.51 mg/l

$$\begin{split} EC_{50} \text{ (48h)} &= 0.20 \text{ mg/l} \\ NOEC &= 0.10 \text{ mg/l} \end{split}$$

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1
Test substance: Santoflex 14 purple liquid, purity: >95%

Remarks: Stock solutions prepared in nanograde Acetone. Water quality

parameters (pH, temperature, dissolved oxygen content) monitored throughout test. Data reported at 95% confidence

level.

Reference: Monsanto ABC 30688, Analytical Bio-Chemistry Labs, 1983

## \*4.3 TOXICITY TO AQUATIC PLANTS, e.g. algae

Species: <u>Selenastrum capricurnutum</u> (freshwater alga)
Endpoint: Biomass [ X ]; Growth rate [ X ]; Other [ ]

Exposure period: 96 Hours

Results:  $EC_{50}$  (24h) = 1.9 ppm

EC50 (96h) = 0.7 ppm NOEC = 0.3 ppm LOEC = 0.6 ppm

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA <u>Selastrum capricornutum</u> Printz Algal Assay Test (1978)

open-system []; closed-system [X]

GLP: Yes [X] No []? [] Klimisch 1
Test substance:. Santoflex 14 reddish purple gel, purity: >95%

Remarks: Stock solutions prepared in reagent grade DMF. Concentrations

determined by range-finding test. Confirmation of effect by <u>in vivo</u> chlorophyll a and cell numbers. Data reported at 95%

confidence level.

Reference: Monsanto BP-81-5-82 EG&G Bionomics, 1981

## 5. TOXICITY

## \*5.1 ACUTE TOXICITY

## 5.1.1 ACUTE ORAL TOXICITY

Type:  $LD_0[]; LD_{100}[]; LD_{50}[X]; LDL_0[]; Other[]$ 

Species/strain: Rats, Sprague-Dawley Albino

Value: 2100 mg/kg b.w.

Discriminating dose: 2510 mg/kg/bw

Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: CP-26658 Lots KC06-14 and KC06-17, purity: >95%

Remarks: Five groups of male and female rats were fed a single oral dose

of the undilated test article via oral gavage. Dosages were 1260, 1580, 2000, 2510 and 3160 mg/kg. Clinical signs of toxicity included reduced activity and appetite for 2-4 days for survivors, and increasing weakness, collapse and death for decedents in 1-4 days. Gross autopsy findings on decedents were hemorragic areas in the lungs, discolored livers and acute gastrointestinal inflammation. Survivors were sacrificed after seven days. All

viscera of survivors appeared normal.

Reference: Monsanto Y-73-169 Younger Laboratories, 1973

#### 5.1.2 ACUTE INHALATION TOXICITY

Type:  $LC_0[\ ];\ LC_{100}[\ ];\ LC_{50}[X];\ LCL_0[\ ];\ Other[\ ]$ 

Species/strain: Rats, Sprague-Dawley Albino

Exposure time: 6 Hours
Value: >0.14 mg/kg
Method: Acute Inhalation

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: CP-26658 liquid, purity: >95%

Remarks: A group of four rats was exposed to the test article at a

concentration of 0.14 mg/l in warm (76.5°F) air for 6 hours. All

animals survived. No clinical signs of toxicity were noted.

Reference: Monsanto Y-67-101, Younger Laboratories, 1967

## 5.1.3 ACUTE DERMAL TOXICITY

Type:  $LD_0[\ ]; LD_{100}[\ ]; LD_{50}[\ X\ ]; LDL_0[\ ]; Other[\ ]$ 

Species/strain: Rabbits, New Zealand Albino

Value: >5010 mg/kg b.w. Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [ X ] Klimisch 2

Test substance: CP-26658 Lots KC06-14 and KC06-17, purity: >95%

Remarks: The undiluted test article was applied to the shaved skin of two

groups of male and female rabbits at dose levels of 5010 and 7940 mg/kg/bw. Clinical signs of toxicity noted were reduced appetite and activity for 4-7 days in survivors, and increased weakness, collapse and death at 8 days for decedents. Gross autopsy findings in decedents included hemorragic areas in the lung, liver and spleen, and discoloration of the kidneys. General gastrointestinal inflammation was also noted. Survivors were

sacrificed after 14 days. All viscera in survivors appeared

normal.

Reference: Monsanto Y-73-169 Younger Laboratories, 1973

## \*5.4 REPEATED DOSE TOXICITY

Species/strain: Rats, Sprague-Dawley Albino

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral/Dietary Exposure period: One Month Frequency of treatment: Daily Post exposure observation period:

Dose: 0, 500, 750, 1500 and 300 ppm Control group: Yes [ **X** ]; No [ ]; No data [ ];

Concurrent no treatment []; Concurrent vehicle[X]; Historical []

NOEL: 500 ppm

LOEL: Not Determined

Results: The test article was administered to groups of 25 male and 25

female rats in a controlled study for one month. Verification of test article stability and dose levels was verified via gas chromatography. Animals were observed twice daily and weighed weekly. Overall averages for dietary concentrations were established as 0, 450, 660, 1300 and 2800 ppm. There were no mortalities during the in-life portion of the study. Toxicity during the in-life phase was indicated by a dose-related reduction of food intake and reduced body weight gains in both males and females at all dietary levels. There were no clinical signs of toxicity observed during the study. There were no gross pathology changes noted at sacrifice which were considered treatment-related, and no significant differences in liver weights or organ coloration. The NOEL for male rats was considered to be 500 ppm. The same NOEL was marginally established for female rats, even though there was a slight, but not statistically

significant difference seen in average body weights.

Method: Dunnett, C.W., A Multiple Comparison Procedure for

Comparing Several Treatments with a Control, Jour. Am. Stat.

Assoc. 50: 1096-1121, 1955

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 14 dark liquid, Lot# KJ08-09, purity: >95% Reference: Monsanto ML-87-309, Environmental Health Lab, 1987

## \*5.5 GENETIC TOXICITY IN VITRO

## A. BACTERIAL TEST

Type: Bacterial Reverse Mutation Assay - Ames

System of testing: Salmonella typhimurium TA-1535 TA-1537 TA-1538 TA-98

TA-100; Saccharomyces cerevisiae D4

Concentration: 0.001, 0.01, 0.1, 1.0 and 5.0 microliters/plate

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 5.0 ul/plate (TA-98 only)

Without metabolic activation: 5.0 ul/plate (TA-98 only)

Precipitation conc: Not Determined

Genotoxic effects: + ?

With metabolic activation: [ ] [ ] [X]
Without metabolic activation: [ ] [ ] [X]

Method: Ames Plate Test (Overlay method) 1975; OECD 471 equivalent

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex 14 dark liquid, purity: >95%

Remarks: The test article, in DMSO solvent, was tested directly and in the

presence of liver microsomal enzyme preparations from Aroclorinduced rats. The test compound did not demonstrate mutagenic activity in any of the assasy conducted and was not considered to

be mutagenic under test conditions.

Reference: Monsanto BIO-76-229, Litton Bionetics, 1976

## B. NON-BACTERIAL IN VITRO TEST

Type: Forward Mutation Mouse Lymphoma Assay System of testing: L5178Y Mouse Lymphoma Cells Concentration: 0.625 - 10.0 nl/ml without activation 1.25 - 60.0 nl/ml with activation Metabolic activation: With []; Without []; With and Without [X]; No data [] Results: Cytotoxicity conc: With metabolic activation: 60 nl/ml Without metabolic activation: 20 nl/ml Precipitation conc: Not Determined Genotoxic effects: + ? -With metabolic activation: [][][X]Without metabolic activation: [][][X] Clive, D., and Spector, J.F.S., Laboratory Procedure for Method: Assessing Specific Locus Mutations at the TK Locus in Cultured L5178Y Mouse Lymphoma Cells. Mutation Res., 31:17-29, 1975 Yes [X] No [] ? [] GLP: Klimisch 1 Santoflex 14 dark liquid, purity: >95% Test substance:. Remarks: The test compound in DMSO solution was evaluated for ability to increase mutations at the TK locus in mouse lymphoma cells at dose ranges of 0.625 to 10 nl/ml without activation and at 1.25 to 60 nl/ml with activation. Dose levels were established during a preliminary range-finding study. The dose levels selected included highly toxic treatments. Even at the highly toxic doses, the mutant frequency was comparable to negative controls. The test substance was considered to be inactive under assay conditions. Reference: Monsanto BO-78-225, Litton Bionetics, 1979 Type: Forward Mutation Assay, CHO/HGPRT System of testing: Chinese Hamster Ovary cells Concentration: 1-10 ug/ml without activation 10-30 ug/ml with activation Metabolic activation: With []; Without []; With and Without [X]; No data [] Results: Cytotoxicity conc: With metabolic activation: 7 ug/ml Without metabolic activation: 5 ug/ml Precipitation conc: Not Determined Genotoxic effects: + ? -With metabolic activation: [][][X] Without metabolic activation: [][][X] Method: CHO/HGPRT Mutation Assay (1981) Hsie, et.al. GLP: Yes [X] No []?[] Klimisch 1 Santoflex 14 liquid Lot# KJ08-09, purity: >95% Test substance:. The mutagenic potential of Santoflex 14 was tested in cultured Remarks: Chinese hamster ovary (CHO) cells. Mutation at the Hypoxanthine guanine phosphororibosyl transferase (HGPRT) locus was measured. Dosages for the test article, dissolved in Acetone, were established with a range-finding experiment. No

Chemical-related mutagenicity was observed in either the initial or the confirmation experiment, with or without S9 activation, were noted. Santoflex 14 was not mutagenic in CHO cells under

any test conditions.

Reference: Monsanto ML-87-340, Environmental Health Labs, 1988

Type: <u>In vitro</u> Cytogenetics Study

System of testing: Chinese Hamster Ovary (CHO) cells

Concentration: 1.5 - 15.0 ug/ml

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 12.5 ug/ml

Without metabolic activation: 12.5 ug/ml

Precipitation conc: Not Determined

Genotoxic effects: + ? -

With metabolic activation: [X] [] [] Without metabolic activation: [X] [] []

Method: Preston, Et. al., Mammalian <u>In vivo</u> and <u>In vitro</u> Cytogenics

Assays: A report to the U.S. Gene-Tox Program (1981)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance:. Santoflex 14 opaque liquid #T870091, purity: >95%
Remarks: Treatment solutions were made using Acetone. Two range-

Finding experiments were run to determine the optimum dose concentrations. MMS and CP were used as concurrent positive controls for treatment with and without S9 activation, respectively. Duplicate samples per treatment condition were used. Scoring for cytogenetic damage was performed on the solvent controls, positive controls, and the three highest dose levels of the test chemical. The cells were scored for both mitotic index and average cell generation time and compared to the

index and average cell generation time and compared to the solvent control. Average cell generation time was 12 hours for both, with a mitotic index of 5-8% Statistically significant increases in number of cells with structural aberrations and average structural aberrations/cell were observed at the 15 ug/ml level for the 48 hour harvest time and for average structural aberrations/cell at the 24 hour harvest time without S9 activation.

aberrations/cell at the 24 hour harvest time without S9 activation. A significant dose-response was not observed. The aberrant cells harvested at 24 and 48 hours included mainly cells with chromatid- and chromosome-type deletions, with a few decentrics and cells with chromatid interchanges. This was also observed in the solvent control. The positive MMS control yielded significant increases in both cells with structural aberrations and number of aberrations/cell. With S9 activation, a statistically significant increase in the number of cells with structural aberrations, and number of aberrations/cell was observed at the 10 ug/ml dose level, and for the number of aberrations/cell at 7.5 ug/ml and 12 hour harvest time. No dose-related response was observed. Aberrations were mainly deletions, with a few cells having

control yielded the expected positive response. A retest confirmed results. Santoflex 14 was concluded to have a weak

chromatid interchanges, intrachanges and triradials. The positive

clastogenicity in CHO cells under test conditions

Reference: Monsanto ML-87-341, Environmental Health Labs, 1989

## \* 5.6 GENETIC TOXICITY IN VIVO

Type: Mammalian Bone Marrow Metaphase Assay

Species/strain: Rats, Sprague-Dawley

Sex: Female []; Male []; Male/Female [X]; No data []

Route of Administration: Oral gavage Exposure period: 6, 18 and 30 hours

Doses: 1100 mg/kg/bw (slightly above 1/4 he oral LD50)

Results:

Effect on mitotic index or P/N ratio:

Genotoxic effects: + ? -

[][][X]

Method: Preston, Et. al., Mammalian <u>In vivo</u> and <u>In vitro</u> Cytogenics

Assays: A report to the U.S. Gene-Tox Program (1981)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex 14 dark oil, purity: >95%

Remarks: Groups of 5 male and female rats were dosed with 1050, 1100,

1200, 1500 and 2000 mg/kg/bw in two range-finding studies. Based upon the results, a dose level of 1100 mg/kg/bw was chosen as close to the maximum tolerated dose for the metaphase analysis. During the In vivo phase, test animals were observed for pharmacotoxicity immediately after dosing, and at 6, 18 and Observations indicated moderate to severe pharmacotoxic signs. Two to three hours prior to sacrifice, each animal received a single intraperitoneal dose of colchicine at 4 mg/kg/bw to arrest dividing cells in metaphase. Both femurs were removed from each animal after sacrifice. The distal end was snipped off one bone and the proximal end off the other. Bone marrow cells were flushed, washed and centrifuged, and slides were prepared using freshly prepared fixative. A total of 500 well-spread metaphase cells with a minimum of overlapping chromosomes were scored for the presence of chromosome aberration per experimental treatment point (50 per animal) by two investigators (25 each per animal). Cells judged acceptable for analysis based on cell morphology and total chromosome number were further analyzed with 100x oil immersion objective where abnormalities were detected and classified. The mean number of aberrations per cell per animal was analyzed for statistically significant increases by one-tailed t tests for each time interval. Santoflex 14 did not produce significant increases in the number of aberrations or in the number of aberrant metaphases at any of the three sacrifice times evaluated. Pharmacotoxic signs observed during the study indicated that the test chemical was dosed near the maximum tolerated dose. Conclusion was that the test chemical was negative in ability to induce structural chromosomal aberrations to the hemopoietic cells of the rat bone marrow under test conditions.

cens of the fat bone marrow under test conditions.

Reference: Monsanto PK-88-342, Pharmakon Research, 1988

*5.8	TOXICITY TO REPRODUCTION					
	Type:	Fertility []; One-generation study []; Two-generation study []; Other [ ]				
	Species/strain:					
	Sex:	Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]				
	Route of Administration	Route of Administration:				
	Exposure period:					
	Frequency of treatmer	nt:				
	- ·	Post exposure observation period:				
	•	Premating exposure period: male: , female:				
	Duration of the test:					
	Doses:	Doses:				
	Control group:	Yes [ ]; No [ ]; No data [ ];				
		Concurrent no treatment []; Concurrent vehicle []; Historical []				
	NOEL Parental:					
	NOEL F1 Offspring:					
	NOEL F2 Offspring:					
	Results:	General parental toxicity:				
		Toxicity to offspring:				
	Method:					
	GLP:	Yes [] No [] ? []				
	Test substance:	, purity:				
	Remarks:					
	Reference:					

## \*5.9 DEVELOPMENTAL TOXICITY/ TERATOGENICITY

Species/strain:

Sex: Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]

Route of Administration:

Duration of the test: Exposure period: Frequency of treatment:

Doses:

Control group: Yes [ ]; No [ ]; No data [ ];

Concurrent no treatment []; Concurrent vehicle []; Historical []

NOEL Maternal Toxicity: NOEL teratogenicity:

Results:

Maternal general toxicity: Pregnancy/litter data:

Foetal data:

Method:

GLP: Yes [ ] No [ ] ? [ ]

Test substance: ..... purity: .....

Remarks: Reference:

#### 5.10 OTHER RELEVANT INFORMATION

## A. Specific toxicities

Type: Immunotoxicity – Repeat Insult Patch Test

Human skin, Santoflex 14 Antiozonant

Shelansky Method (Procedings of the Toilet Goods

Association, No. 19, May 1953)

Results: Fifty human volunteers not previously exposed to test rubber

formulations were selected. Squares soaked in the test material were applied to the arm or back and held in place with tape. Patches were removed after 24 hours and the sites examined for reactions, after which the material was reapplied. Fifteen such primary applications were made, followed by a 2-week rest period. A challenge application was then applied as before, and to the same site. No reactions were produced by either the primary or challenge applications. There was no evidence of primary irritation or skin fatigue. There was no evidence of skin

sensitization under the test conditions.

Remarks: Concentration of test article was not noted. Both male and female

volunteers were used in the study.

Reference: Monsanto SH-65-3, Industrial Biology Labs, 1965

Type: Immunotoxicity – Repeat Insult Patch Test

Human skin, Unvulcanized Rubber containing Santoflex 14

Antiozonant

Shelansky Method (Procedings of the Toilet Goods

Association, No. 19, May 1953)

Results: Fifty one human volunteers not previously exposed to test rubber

formulations were selected. The test material, in the form of 1"

squares of unvulcanized rubber, was affixed to the upper arm of each test subject and covered with gauze (occluded). Patches were removed after 24 hours and the sites examined for reactions. Direct effects by single contact were graded with a numerical score ranging from 0 (no response) to 4 (severe response) for primary irritation. Choice of contact site for the second and all subsequent applications was based on the condition of the skin at the original contact site. If irritation occurred, a different site was chosen. If no irritation occurred, the test patch was reapplied to the same site. There were 15 such applications in the induction phase of the study. Following a 14-day rest period, a challenge application was applied at the original contact site. No visible skin changes were noted on any test subject during

negative for delayed contact hypersensitivity.

Remarks: Concentration of test article in the rubber compound was 3 parts

per 100 parts of SBR 1000 rubber (3 phr) Both males and

either the induction phase or the challenge phase of the study. The test article was considered to be negative for primary skin irritation, negative for skin fatigue by sequential contact, and

females were used in the study.

Reference: Monsanto SH-67-13, Industrial Biology Labs, 1967

## B. Toxicodynamics, toxicokinetics

Type: Results: Remarks:

References:

# \* 5.11 EXPERIENCE WITH HUMAN EXPOSURE Results: .

Remarks:
Reference:

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IUCLID

Data Set

Existing Chemical ID: 15233-47-3 CAS No. 15233-47-3

TSCA Name 1,4-benzenediamine, N-(1-methylheptyl)-N'-phenyl-

EINECS No. 239-281-1

Molecular Weight 296

Producer Related Part

Company:

Creation date: 08-NOV-2001

Substance Related Part

Company:

Creation date: 08-NOV-2001

Memo: RAPA PPD Category

Printing date: 09-NOV-2001

Revision date:

Date of last Update: 09-NOV-2001

Number of Pages: 19

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 09-NOV-2001

1. General Information ID: 15233-47-3

1. General information 1D. 15255-47-5

## 1.0.1 OECD and Company Information

Type: lead organisation

Name: American Chemistry Council (formerly Chemical Manufacturers

Association) Rubber and Plastics Additives (RAPA) HPV Panel

Street: 1300 Wilson Boulevard Town: 22209 Arlington, VA

Country: United States
Phone: 703-741-5600
Telefax: 703-741-6091

08-NOV-2001

Type: cooperating company
Name: Bayer Corporation
Country: United States

08-NOV-2001

Type: cooperating company

Name: Ciba Specialty Chemicals Corporation

Country: United States

08-NOV-2001

Type: cooperating company Name: Crompton Corporation

Country: United States

08-NOV-2001

Type: cooperating company Name: Flexsys America L.P.

Country: United States

08-NOV-2001

Type: cooperating company

Name: Noveon, Inc (formerly BF Goodrich)

Country: United States

08-NOV-2001

Type: cooperating company

Name: R.T. Vanderbilt Company, Inc.

Country: United States

08-NOV-2001

Type: cooperating company

Name: The Goodyear Tire & Rubber Company

Country: United States

08-NOV-2001

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Date: 09-NOV-2001

1. General Information

Date: 15233-47-3

Type: cooperating company
Name: The Lubrizol Corporation

Country: United States

08-NOV-2001

Type: cooperating company

Name: UOP, LLC. Country: United States

08-NOV-2001

1.0.2 Location of Production Site

-

1.0.3 Identity of Recipients

\_

1.1 General Substance Information

Substance type: organic Physical status: liquid Purity: > 95 % w/w

08-NOV-2001

1.1.0 Details on Template

\_

1.1.1 Spectra

-

## 1.2 Synonyms

N-phenyl - N'-(1-methylhepyl)-p-phenylenediamine 08-NOV-2001

UOP 688 Antiozonant 08-NOV-2001

1.3 Impurities

\_

1.4 Additives

\_

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Date: 09-NOV-2001

1. General Information ID: 15233-47-3

1.5 Quantity

-

1.6.1 Labelling

-

1.6.2 Classification

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1.7 Use Pattern

-

1.7.1 Technology Production/Use

\_

1.8 Occupational Exposure Limit Values

\_

1.9 Source of Exposure

-

1.10.1 Recommendations/Precautionary Measures

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1.10.2 Emergency Measures

\_

1.11 Packaging

-

1.12 Possib. of Rendering Subst. Harmless

-

1.13 Statements Concerning Waste

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1.14.1 Water Pollution

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1.14.2 Major Accident Hazards

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Date: 09-NOV-2001

1. General Information

ID: 15233-47-3

1.14.3 Air Pollution

-

1.15 Additional Remarks

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1.16 Last Literature Search

-

1.17 Reviews

-

1.18 Listings e.g. Chemical Inventories

-

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## 2.1 Melting Point

Value:

Remark: Unknown, no studies available

08-NOV-2001

## 2.2 Boiling Point

Value: 431 degree C at 1013 hPa

Method: other: no data

GLP: no

08-NOV-2001 (1)

## 2.3 Density

Type: relative density
Value: 1.003 at 15.6 degree C

Method: other: no data

GLP: no

Result: Specific gravity = 1.003

08-NOV-2001 (1)

## 2.3.1 Granulometry

-

## 2.4 Vapour Pressure

Value:

Remark: Unknown, no studies available

08-NOV-2001

## 2.5 Partition Coefficient

log Pow:

Method: OECD Guide-line 107 "Partition Coefficient (n-octanol/water),

Flask-shaking Method"

Year:

Result: Method not applicable.

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

08-NOV-2001 (2)

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## 2.6.1 Water Solubility

Qualitative: not soluble

Method: OECD Guide-line 105 "Water Solubility"

Remark: Evaluation as part of Certificate of Analysis

Result: Insoluble;

pH Value, pKa Value: Unknown, no studies available

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

08-NOV-2001 (2)

## 2.6.2 Surface Tension

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#### 2.7 Flash Point

\_

## 2.8 Auto Flammability

\_

## 2.9 Flammability

\_

## 2.10 Explosive Properties

-

## 2.11 Oxidizing Properties

Result:

Remark: Unknown, no studies available

08-NOV-2001

## 2.12 Additional Remarks

Memo: Fat Solubility

Method: OECD 116 Result: 100%

08-NOV-2001 (2)

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## 3.1.1 Photodegradation

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 1560000 molecule/cm3

Rate constant: .000000000229 cm3/(molecule \* sec)

Degradation: 50 % after .6 hour(s)

Method: other (calculated): AOP Program (v1.89) 1999 Year:

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Acceted calculation method

Flag: Critical study for SIDS endpoint

08-NOV-2001 (3)

## 3.1.2 Stability in Water

## 3.1.3 Stability in Soil

## 3.2 Monitoring Data (Environment)

## 3.3.1 Transport between Environmental Compartments

Type: fugacity model level III

Media: other: air - water - soil - sediment

Air (Level I): Water (Level I): Soil (Level I): Biota (L.II/III): Soil (L.II/III):

Method: other: EPIWIN, Level III Fugacity Model

1999 Year:

Result: Media Concentration Half-Life Emissions Fugacity

	(percent)	(hr)	(kg/hr)	(atm)
Air	0.0248	1.12	1000	7.34e-013
Water	8.94	900	1000	2.61e-014
Soil	43.4	900	1000	3.56e-016
Sedimen	t 47.6	3.6e+003	0	1.76e-014

Media	Reaction	Advection	Reaction	Advection
	(kg/hr)	(kg/hr)	(percent)	(percent)
Air	615	9.94	20.5	0.331
Water	275	358	9.18	11.9
Soil	1.34e+003	0	44.6	0
Sediment	367	38.1	12.2	1.27

Persistence Time: 1.33e+003 hr

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Date: 09-NOV-2001
3. Environmental Fate and Pathways

ID: 15233-47-3

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Reaction Time: 1.54e+003 hr Advection Time: 9.86e+003 hr

Percent Reacted: 86.5
Percent Advected: 13.5
Reliability: (2) valid with restrictions
Acceted calculation method

Flag: Critical study for SIDS endpoint

08-NOV-2001 (3)

3.3.2 Distribution

-

3.4 Mode of Degradation in Actual Use

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3.5 Biodegradation

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3.6 BOD5, COD or BOD5/COD Ratio

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3.7 Bioaccumulation

-

3.8 Additional Remarks

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#### AQUATIC ORGANISMS

## 4.1 Acute/Prolonged Toxicity to Fish

Type: other

Species: other: Freshwater fish

Exposure period: 96 hour(s)

mg/lAnalytical monitoring: no

LC50: .067

Method: other: ECOSAR Program (v0.99e)

1999 GLP: no Year:

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Acceted calculation method

Critical study for SIDS endpoint Flag:

08-NOV-2001 (3)

other Type:

other: Saltwater fish Species:

Exposure period: 96 hour(s)

Analytical monitoring: no Unit: mq/1

LC50: .094

Method: other: ECOSAR Program (v0.99e)

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Acceted calculation method

Critical study for SIDS endpoint Flaq:

08-NOV-2001 (3)

## 4.2 Acute Toxicity to Aquatic Invertebrates

Type: other

Species: Daphnia sp. (Crustacea)

Exposure period: 48 hour(s)

mg/lUnit: Analytical monitoring: no

LC50 : .093

Method: other: ECOSAR Program (v0.99e)

1999 Year: GLP: no

other TS: molecular structure Test substance:

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Acceted calculation method

Flaq: Critical study for SIDS endpoint

08-NOV-2001 (3)

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Date: 09-NOV-2001 ID: 15233-47-3 4. Ecotoxicity

Type: other

Species: Mysidopsis bahia (Crustacea)

Exposure period: 96 hour(s)

Unit: mg/1Analytical monitoring: no

LC50 : .00134

Method: other: ECOSAR Program (v0.99e)

1999 GLP: no Year:

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Acceted calculation method

(3) 08-NOV-2001

## 4.3 Toxicity to Aquatic Plants e.g. Algae

Species: other algae: Green algae

Endpoint:

Exposure period: 96 hour(s)

Unit: mq/1Analytical monitoring: no

.072 EC50:

other: ECOSAR Program (v0.99e) Method:

Year: 1999 GLP: no

Test substance: other TS: molecular structure

Remark: Chemical may not be soluble enough to measure this predicted

effect.

Reliability: (2) valid with restrictions

Acceted calculation method

Flag: Critical study for SIDS endpoint

08-NOV-2001 (3)

## 4.4 Toxicity to Microorganisms e.g. Bacteria

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4. ECOCOXICICY

- 4.5 Chronic Toxicity to Aquatic Organisms
- 4.5.1 Chronic Toxicity to Fish

-

4.5.2 Chronic Toxicity to Aquatic Invertebrates

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## TERRESTRIAL ORGANISMS

4.6.1 Toxicity to Soil Dwelling Organisms

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4.6.2 Toxicity to Terrestrial Plants

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4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

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4.7 Biological Effects Monitoring

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4.8 Biotransformation and Kinetics

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4.9 Additional Remarks

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## 5.1 Acute Toxicity

## 5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain: other: Holtzman

Sex: male

Number of

Animals: 5

Vehicle: other: corn oil Value: 4.3 mg/kg bw

Method: other: Method described by Weil, C.S., Biometrics 8, 249, 1952

Year: 1952 GLP: no

Test substance: other TS: Commercial product, >95% purity

Method: UOP 688 was administered orally to six groups , each composed

of 5 male albino rats, weight range 219-251 grams. Each dose was administered either undiluted or as a 10% volume/volume solution in corn (Mazola) oil. Dosage levels tested were 0.046, 0.10, 2.15, 4.46, 10.0, and 21.5 mg/kg body weight. All animals were observed closely for gross signs of systemic toxicity and mortality during the day of dosage, and at least once daily thereafter for 14 days. All animals were subject

to gross necropsy at study termination.

Result: Animals in the 0.046, 0.1, and 2.15 mg/kg dosage levels

generally exhibited normal appearance and behaviour throughout the 14 day period. Rats at the 4.64 mg/kg dose level began showing depression, slowed righting reflexes, and diarrhea on the second day following dosage. On the fourth day after dosage, one rat showed labored respiration, ataxia, depressed righting, placement, and pain reflexes, and a marked bloody nasal discharge. These signs generally continued until death occurred, or until the fifth day following dosage when the two surviving rats appeared normal. The rats in the 10.0 and 21.5 mg/kg doe levels showed diarrhea, unkempt fur, depression, depressed relexes, and a dark oily stain in the perineal area on the day after dosage. These signs continued until death

occurred. Death was preceded by lacrimation and coma.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

08-NOV-2001 (4)

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## 5.1.2 Acute Inhalation Toxicity

Type: Species: Strain: Sex: Number of Animals: Vehicle: Exposure time:

Value: Method:

Year: GLP:

Test substance:

Unknown, no studies available.

Not an appropriate route of exposure due high boiling point.

08-NOV-2001

## 5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain: New Zealand white Sex: male/female

Number of

10 Animals:

Vehicle:

Value: > 2000 mg/kg bw

Method: other: U.S. Code of Federal Regulations 40 CFR 163

Year: GI.P:

other TS: Commercial product, >95% purity Test substance:

Method: The test material was applied to five male and five female

white New Zealand white rabbits. The dose was applied to the abdominal skin which had been previously been shaven. The abdominal skin area of all the rabbits was abraded by making a series of longitudinal minor epidermal incisions placed two to

three centimeters apart, using a hypodermic needle as a cutting tool. The abrasions were sufficiently deep to penetrate the epidermis, but not to induce bleeding. The undiluted sample was applied at a dosage level of 2.0 grams/kg of body weight. The test sample was kept in contact with the skin on at least 10% of the body surface. During the exposure period, each rabbit was observed for signs of toxicity at two, four and five and one half hours post application. After 23 1/4 to 24 hours of skin contact exposure, any unabsorbed sample remaining on the skin was removed by gentle sponging with a moistened towel. Rabbits were observed for 14 days following completion of the exposure period. Examinations for gross signs of systemic toxicity were carried out twice daily during

rabbits were weighted, sacrificed and gross necropsy was

this period. At the end of the 14 day observation period,

performed.

study reviewed by lab QA Director Remark:

Result: One female rabbit was found dead on day two. Necropsy

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revealed diarrhea stains around the anus, congested lungs, a mottled and darkened liver, stomach and intestine which appeared autolytic and pale but congested kidneys. Erythemia

and edema followed by desquamation and atonia were seen at the application site in all surviving animals. Four rabbits exhibited spotted whitening on the day of exposure completion. Systemic effects were limited to transient nasal discharge in

Systemic effects were limited to transient nasal discharge in two animals and transient green colored urine in one animal.

Reliability: (1) valid without restriction

Meets National standards method

Flag: Critical study for SIDS endpoint

08-NOV-2001 (5)

5.1.4 Acute Toxicity, other Routes

-

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration:

Exposure: Semiocclusive

Exposure Time: 24 hour(s)

Number of

Animals: 6
PDII: 1.5

Result:

EC classificat.:

Method: other: U.S. Code of Federal Regulations 40 CRF 163

Year: GLP:

Test substance: other TS: Commercial product, >95% purity

Method: 0.5 ml undiluted test material was applied under one inch

square surgical gauze patches to two abraded skin areas and two intact skin areas on each of six New Zealand White

rabbits. After 24 hours of skin contact exposure, any

unabsorbed sample remaining on the skin was removed by gentle sponging with a moistened towel. The reactions were scored immediately after removal of the patches (24 hour reading),

and again two days later (72 hour reading).

Remark: study reviewed by lab QA Director

Result: Irritative effects noted during the course of the study

included very slight to well defined erythema, at the abraded and intact sites of all animals. Very slight to slight edema scores were noted in five animals on the abraded and intact sites. The Primary Irritation Index was found to be 1.5. Some loss of skin resiliency (atonicity) was noted. No

evidence of corrosivity was observed.

Reliability: (1) valid without restriction

Meets National standards method

09-NOV-2001 (5)

Species: rabbit

Concentration: undiluted

Exposure: Semiocclusive

Exposure Time:

Number of

Animals: 6

PDII: Result:

EC classificat.:

Method: other: U.S. Code of Federal Regulations 49 CFR 173.136 -137

Year: 1992 GLP: yes

Test substance: other TS: Commercial product, Lot #0483, >95% purity

Method: The primary dermal irritation/corrosivity potential was

evaluated when applied to the skin of 3 male and 3 female rabbits under 3 minute, 1 hour, and 4 hour semi-occluded conditions. Each application site was examined for erythemia

and edema according to the Draize method.

Result: No evidence of corrosion was observed at any of the test sites

for any of the exposure periods.

Not considered corrosive to the skin of rabbits

Reliability: (1) valid without restriction

GLP Guideline study

09-NOV-2001 (6)

#### 5.2.2 Eye Irritation

Species: rabbit

Concentration: undiluted

Dose: .1 ml

Exposure Time:

Comment: other: see method

Number of Animals: 9

Result:

EC classificat.:

Method: other: U.S. Code of Federal Regulations 40 CFR 163

Year: GLP:

Test substance: other TS: Commercial product, >95% purity

Method: 0.1 ml of the undiluted test material was applied to the left or right eye of each of nine rabbits. The opposite eye served

as a control. The treated eyes of six rabbits were left unrinsed. The treated eye of three rabbits were rinsed after 30 seconds for 60 seconds with 200 ml of lukewarm water. Examinations for gross signs of eye irritation were made approximately 24, 43, and 70 ½ hours and four, seven, ten, thirteen, sixteen, and nineteen days following application.

Scoring of irritative effects was according to the method of

Draize.

Remark: study reviewed by lab QA Director

Result: Non-rinsed eyes - Irritative effects noted during the study

included isolated occurrences of mild corneal opacity with up

to one-quarter of the corneal area involved in the two

- 15/19 -

rabbits. Conjuctival effects included isolated occurrences of mild erythema in five rabbits. Total irritation score ranged from 0-5.

Rinsed eyes - Mild corneal irritation was observed in the rinsed eye group. These effects generally cleared after four days post-treatment with opacity occurring once after this reading in one rabbit. Sporadic occurrences of mild to moderate conjunctival irritation on days 13 and 19 were noted in three rabbits. The total irritation scores ranged from 0-7.

09-NOV-2001 (5)

#### 5.3 Sensitization

Type: Patch-Test Species: human

Number of

Animals: 15

Vehicle: other: acetone Result: not sensitizing Classification: not sensitizing

Method: other: Adapted from the repeated insult patch test procedure

described by Draize (Appraisal of the Safety of Chemicals in Foods, Drugs, and Cosmetics, pp. 52-55, The Association of

Food and Drug Officials of the United States, 1959)

Year: GLP: no

Test substance: other TS: Commercial product

Method: 0.1 ml o

0.1 ml of a 20% acetone solution of the sample (equivalent to 20 mg of the test material) was applied to a  $\frac{3}{4}$  x  $\frac{7}{8}$  inch piece of filter paper. After the acetone had evaporated, the filter paper was place on the skin of 15 human subjects. Nine patch applications were made to the same location on the upper arm over a period of two weeks. A challenge patch was applied

to skin area not previously exposed to the test material. None of the 15 subjects tested exhibited any evidence of

sensitization.

09-NOV-2001 (7)

5.4 Repeated Dose Toxicity

-

Result:

Date: 09-NOV-2001
5. Toxicity ID: 15233-47-3

# 5.5 Genetic Toxicity 'in Vitro'

Type: Ames test

System of

testing: Salmonella typhimurium strains TA-1535, TA-1537, TA-1538,

TA-98, and TA-100

Concentration: 0.0005, 0/001, 0.0025, 0.005, 0.01, 0.05, 0.1, 0.5 ug/plate Cytotoxic Conc.: Without metabolic activation: >0.07 ug/plate; Precipitation

conc: 0.59 ug/plate

Metabolic

activation: with and without

Result: negative

Method: other: Ames Salmonella/Microsome Plate Test, Protocol 401,

Edition 14

Year: GLP: yes

Test substance: other TS: Commercial product, purity >95%

Remark: Examination of mutagenic activity in the presence and absence

of liver microsomal preparations was conducted. Solvent control (dimethyl sulfoxide) and specific positive control compounds were assayed concurrently with the test material. The concurrent solvent control data were used as a basis for

evaluating results.

Result: The test material did not exhibit genetic activity in any of

the assays conducted and was not mutagenic to the S.

typhimurium indicator organism under the test conditions.

Reliability: (1) valid without restriction

GLP Guideline study

Flag: Critical study for SIDS endpoint

09-NOV-2001 (8)

5.6 Genetic Toxicity 'in Vivo'

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5.7 Carcinogenicity

\_

5.8 Toxicity to Reproduction

-

5.9 Developmental Toxicity/Teratogenicity

\_

5.10 Other Relevant Information

\_

5.11 Experience with Human Exposure

-

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Date: 09-NOV-2001
6. References ID: 15233-47-3

20 20 20 27 2

- (1) From internal technical bulletin, 1981
- (2) Evaluation as part of Certificate of Analysis, by Fine Pharmaceutical Laboratories, Ltd., Hamilton, Ontario, Canada, January 24, 2001
- (3) Meylan W. and Howard P. (1999) EPIWin Modeling Program. Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.
- (4) Unpublished study, "Acute Oral Adminstration of UOP 604 and UOP 688 to Rats", Hill Top Research Institute, Inc. Miamiville, OH, February 13, 1963
- (5) Unpublished study, "Acute Dermal Toxicity, Primary Skin Irritation and Acute Eye Irritation Potential of UOP 688", Hill Top Research, Inc., Cincinnati, OH, September 22, 1981
- (6) Unpublished study, "Skin Corrosivity Study of UOP 688 in Rabbits (DOT/UN Regulations)", Hazelton Wisconsin, Inc, Madison WI, June 25, 1993.
- (7) Unpublished study, "Repeated Insult Patch Test of UOP 688 and 12267", Hill Top Research, Inc., September 20, 1962.
- (8) Unpublished study, "Mutagenicity Test on XPA-28-86/UOP 688 in the Ames Salmonella/Micorsomal Reverse Mutation Assay", Hazelton Laboratories America, Inc., Kensington, MD, October 13, 1981.

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7. Risk Assessment Date: 09-NOV-2001 ID: 15233-47-3

7.1 End Point Summary

-

7.2 Hazard Summary

-

7.3 Risk Assessment

-

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IUCLID

Data Set

Existing Chemical ID: 68953-84-4 CAS No. 68953-84-4

EINECS Name N,N'-diaryl-p-phenylenediamines

EINECS No. 273-227-8

Producer Related Part

Company: Goodyear Chemicals Europe

Creation date: 06-APR-1998

Substance Related Part

Company: Goodyear Chemicals Europe

Creation date: 06-APR-1998

Printing date: 30-OCT-2001

Revision date:

Date of last Update: 20-FEB-2001

Number of Pages: 28

Chapter (profile): Chapter: 2.1, 2.2, 2.4, 2.5, 2.6.1, 3.1.1, 3.1.2, 3.3.1,

 $3.5,\ 4.1,\ 4.2,\ 4.3,\ 5.1.1,\ 5.1.2,\ 5.1.3,\ 5.1.4,\ 5.4,\ 5.5,$ 

5.6, 5.8, 5.9

Reliability (profile): Reliability: 1, 2

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 30-OCT-2001
2. Physico-chemical Data

ID: 68953-84-4

### 2.1 Melting Point

Value: 90 - 105 degree C

Decomposition: ambiguous

Method: other: ASTM D-1519

Year: 1993 GLP: no

Reliability: (2) valid with restrictions

Although this study was probably not conducted to GLP, the

test parameters used were based on a known and well

established procedure.

31-JUL-2000 (34)

#### 2.2 Boiling Point

\_

#### 2.4 Vapour Pressure

-

#### 2.5 Partition Coefficient

log Pow: 3.4 - 4.3

Method: OECD Guide-line 117 "Partition Coefficient (n-octanol/water),

HPLC Method"

Year: 1995 GLP: yes

Remark: The product exhibits much lower values than DDT (6.2) which

provides a benchmark for highly bioaccumulative chemicals.

The test substance contains 3 major components.

Result: # Methyl Groups -0 log Pow 3.37 # Methyl Groups -1 log Pow 3.82

# Methyl Groups -1 log Pow 3.82
# Methyl Groups -2 log Pow 4.28

The major components of the test substance displayed partion

coefficients between 3.4 and 4.3.

Reliability: (1) valid without restriction

01-AUG-2000 (28)

log Pow: > 3.7 at 22.8 degree C

Method: other (measured)

Year: 1992 GLP: yes

Remark: for N,N'-Diphenyl-p-phenylenediamine

Reliability: (1) valid without restriction

20-FEB-2001 (9)

- 1/28 -

log Pow: > 4.3 at 22.8 degree C

Method: other (measured)

Year: 1992 GLP: yes

Remark: For N-phenyl-N'-(o-tolyl)-p-phenylenediamine

Reliability: (1) valid without restriction

31-JUL-2000 (9)

log Pow: > 4.6 at 22.8 degree C

Method: other (measured)

Year: 1992 GLP: yes

Remark: For N,N'-Di(o-tolyl)-p-phenylenediamine

Reliability: (1) valid without restriction

20-FEB-2001 (9)

2.6.1 Water Solubility

-

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Date: 30-OCT-2001
3. Environmental Fate and Pathways ID: 68953-84-4

#### 3.1.1 Photodegradation

-

#### 3.1.2 Stability in Water

Type: Method:

Year: 1994 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: See Biodegradation Studies

Reliability: (1) valid without restriction

31-JUL-2000

#### 3.3.1 Transport between Environmental Compartments

\_

#### 3.5 Biodegradation

Type: anaerobic

Inoculum: activated sludge, domestic

Concentration: 100 mg/l related to Test substance

Degradation: .64 % after 28 day

Result: other: not readily biodegradable

Method: OECD Guide-line 301 F "Ready Biodegradability: Manometric

Respirometry Test"

Year: 1994 GLP: yes

Test substance: as prescribed by 1.1 - 1.4 Reliability: (1) valid without restriction

31-JUL-2000 (22)

Type: anaerobic

Inoculum: activated sludge Degradation: 0 % after 28 day

Method: other: OECD 301 Manometric Respirometry, modified according to

EEC Round RobinTest "Assessment of Respirometry" DGX 1/283/82

Rev. 6, EEC Directive 79/831, Annex V, Part C

Year: 1990 GLP: yes

Test substance: as prescribed by 1.1 - 1.4
Reliability: (1) valid without restriction

31-JUL-2000 (6)

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Date: 30-OCT-2001
4. Ecotoxicity ID: 68953-84-4

AQUATIC ORGANISMS

# 4.1 Acute/Prolonged Toxicity to Fish

Type: flow through

Species: Cyprinus carpio (Fish, fresh water)

Exposure period: 14 day

Unit: mg/l Analytical monitoring: yes

NOEC: .28 LC50: .43

Method: OECD Guide-line 204 "Fish, Prolonged Toxicity Test: 14-day

Study"

Year: 1996 GLP: yes

Test substance: other TS

Method: Test water was generated by adding the test substance in

acetone to a larger volume of water which was stirred, allowed to settle, and then siphoned to a stock solution holding tank. This stock solution was then metered into exposure tanks for the fish experiments. A range-finding trial exposed carp to to nominal levels of 2.5, 5, 10, and 25 mg/L (ppm) of the test substance. Survival rates were up to 80% within the first 48 hours for the three (3) highest dose levels and the 2.5 mg/L induced no mortality in the first 48 hours although 90% deaths were seen through Day six

(6).

In the definitive phase, duplicate test tanks contained 10 carp each and the test substance nominal concentrations of 0, 0.1, 0.23, 0.51, 1.1, and 2.5 mg/L (ppm). Chemical analysis (HPLC) of the test substance in the test tanks on Days -0, -3, -7, and -14 showed that mean concentrations for the 14-day test period were 0.053, 0.12, 0.19, 0.28, and

 $0.67~{
m mg/L}~({
m ppm})$ . Fish densities were  $0.35~{
m g}$  biomass/L flowing test solution per day. Tank volume turnover for the flow-through system was  $6.5/{
m day}$ . Carp were monitored daily for mortality and signs of erratic swimming behavior for 14 days during exposure. Body weights and lengths were recorded for representative fish prior to study initation, and on all

test fish on Day 14. A LC50 value was then calculated.

Result: Carp died only at the highest test substance concentration;

2/20 on Day-3, 7/20 on Day-7, and 20/20 by Day-14. Other findings at the 0.67 mg/L (ppm) level included darkened pigmentation on the fish (likely due to adsorption of the test chemical), lethargic swimming behavior, and loss of equilibrum. There were no test substance-related effects on

body lengths or weights.

Test substance: Tested as the commercial product Reliability: (1) valid without restriction

20-FEB-2001 (29)

- 4/28 -

Date: 30-OCT-2001
4. Ecotoxicity ID: 68953-84-4

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Type: flow through

Species: Oncorhynchus mykiss (Fish, fresh water)

Exposure period: 14 day

Unit: mg/l Analytical monitoring: yes

NOEC: .14 LC50: .26

Method: OECD Guide-line 204 "Fish, Prolonged Toxicity Test: 14-day

Study"

Year: 1997 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: Test water was generated by adding the test substance in

acetone to a larger volume of water which was stirred, allowed to settle, and then siphoned to a stock solution holding tank. This stock solution was then metered into exposure tanks for fish experiments. A preliminary study in trout was performed using nominal concentrations of the test substance of 0.1, 0.23, 0.51, 1.1, and 2.5 mg/L. Mortality rates were 100% at the highest level by Day-3, and was 80%

by Day-7 at 1.1 mg/L.

In the definitive phase, duplicate test tanks contained 10 trout each, Test substance nominal concentrations of 0, 0.094, 0.19, 0.38, 0.75, and 1.5 mg/L (ppm) were chosen. Chemical analysis (HPLC) of the test substance in the test tanks on Days -0, -7 and -14 showed that mean concentrations for the 14-day test period were 0.062, 0.093, 0.14, 0.35, and 0.66 mg/L (ppm). Fish densities were 0.079 g biomass/L

flowing test solution per day. Tank volume turnover for the flow-through system was  $6.5/\mathrm{day}$ . Fish were monitored daily for mortality and signs of erratic swimming behavior for 14-days during exposure. Body weights and lengths were recorded for representative fish prior to study initiation, and on all test fish on Day-14. LC50 values were calculated

for 96-hours and 14-days.

Result: Fish died only at 0.35 and 0.66 mg/L concentrations; 0/20

and 1/20 died by Day-2 and 1/20 and 19/20 by Day -4, respectively. Further, 100 % of the high dose (0.66 mg/L) fish died by Day-5 and 17/20 of the 0.37 mg/L fish by Day-14. Other findings at the two highest levels included darkened pigmentation of the fish, lethargic swimming behavior, and loss of equilibrium. There were test

substance-related effects on 14-day body lengths and weights in the 0.35 mg/L group. The calculated LC50 for the test substance in the study at 96-hours was 0.48 mg/L and 0.26 mg/L at 14-days. The No Observed Effect Concentration (NOEC)

was 0.14 mg/L at 96-hours and 14-days.

Reliability: (1) valid without restriction

31-JUL-2000 (37)

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Date: 30-OCT-2001
4. Ecotoxicity ID: 68953-84-4

#### 4.2 Acute Toxicity to Aquatic Invertebrates

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: yes

NOEC: .36 EC50: 1.8

Method: OECD Guide-line 202, part 1 "Daphnia sp., Acute

Immobilisation Test"

Year: 1996 GLP: yes

Test substance: other TS

Method: A range-finding study used ten (10) 24-hour old daphnids

exposed to nominal levels of 0, 13,22,36,60, and 100 mg/L of the test substance. Immpbilization (15%) of the daphnids occurred at the highest level (100 mg/L). Sublethal lethargy was observed at all but the lowest test concentration (13 mg/L). Brown matter, apparently the test substance since brown precipitate was observed in the media, was observed to

adhere to both surviving and non-surviving daphnids.

In the definitive phase, duplicate aquaria containing 10

daphnids each and test substance nominal concentrations of 0, 1.3, 2.2, 3.6, 6.0 and 10  $\ensuremath{\text{mg/L}}$  (ppm) were prepared. Mean values for the test substance concentrations in the test media were determined by averaging chemical analyses (HLPC) of 0-hours and 48-hours.

Daphnia immobilization and aquaria observations were made at 24- and 48-hours following the study initation. From these data, an Effective Concentration in one-half the organisims (EC50) and a No Observed Effect Concentration (NOEC) were estimated.

Result:

Measured concentrations of the test substance ranged from 19 to 29% of nominal levels. At the highest concentration (1.8 mg/L), 25 % of the daphnids were immobilized at 48-hours of exposure. For the 0.68 and 1.1 mg/L groups, Five (5) % of the daphnids were immobile. No immobilization was observed at 0.20 and 0.36 mg/L exposures. Lethargic activity was not observed at any treatment level. Brown particulates, perhaps the test substance, were observed to adhere to the test daphnids, with some buoyed to the surface of the aquaria by this particulate material. The results indicated that the EC50 for the test substance was 1.8 mg/L. The No Observed Effect Concentration (NOEC) was shown to be 0.36 mg/L. Tested as the commercial product

Test substance: Reliability:

(1) valid without restriction

31-JUL-2000 (27)

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4. Ecotoxicity

Date: 30-OCT-2001 ID: 68953-84-4

#### 4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Selenastrum capricornutum (Algae)

Endpoint: biomass Exposure period: 72 hour(s)

Unit: μg/l Analytical monitoring: yes

NOEC: 4.3 EC10: 4.3 EC50: 18

OECD Guide-line 201 "Algae, Growth Inhibition Test" Method: Year: 1996

other TS Test substance:

Method: A range-finding trial used nominal levels of 0, 1,10, 100,

> and 1000 ug/L (ppb) of the test substance and a solvent control in algae cultures (approximately 1x104 cells per flask). Following 72-hours incubation, algal cell densities

were determined using a hemacytometer. Values were

127,76,109,69 and 1%, respectively, of the solvent control response. These values were used to set exposures for the definitive phase.

In the definitive phase, triplicate algal cultures were exposed to the test substance at nominal concentrations of 16, 31, 63,130, 250, and 500 ug/L (ppb). Cell densities were monitored at 24-, 48-, and 72-hours following study initiation. From these data, EC50 (50% decrease) values for Biomass (EbC50) and Growth Rate (ErC50) were calculated. Test substance concentrations in the test media were determined at 0- and 72-hours using HLPC. The mean concentrations were 7.5, 13, 14, 28, 50, and 79 ug/L (ppb). The inhibitions of algae Growth Rates for the test substance in the definitive 72-hour study were 0, 2, 15, 20, 32, and 38% (relative to pooled control values) for the measured test substance concentrations of 7.5, 13, 14, 28, 50, and 79 ug/L (ppb). Corresponding inhibitions of Biomass generation were 15, 41, 59, 63, 81, and 91%. Individual cell appearances were found microscopically to be normal for surviving cells except cellular bloating was noted at the

highest exposure level. Calculations indicated that the ErC50 for the test substance was > 79 ug/L (ppb) while the EbC50 was 18 ug/L (ppb). The No Observed Effect

Concentrations (NOECs) were assumed to be equivalent to EC10 values, and accordingly were EbC10 = 4.3 ug/L (ppb) and

ErC10 = 31 ug/L (ppb).

The EC50 values for the test substance ranged from 18 to > 79 ug/l (ppb) for Biomass increases and Growth Rates. The NOECs ranged from 4.3 to 31 ug/L (ppb) for these parameters.

Test substance: Tested as the commercial product Reliability:

Result:

(1) valid without restriction

31-JUL-2000 (30)

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Date: 30-OCT-2001 ID: 68953-84-4 4. Ecotoxicity

Species: Selenastrum capricornutum (Algae)

Endpoint: growth rate Exposure period: 72 hour(s)

Unit: μg/l Analytical monitoring: yes

NOEC: 31 EC10: 31 EC50: > 79

Method: OECD Guide-line 201 "Algae, Growth Inhibition Test" Year: 1996 GLP: yes

Test substance: other TS

Method: A range-finding trial used nominal levels of 0, 1,10, 100,

and 1000 ug/L (ppb) of the test substance and a solvent

control in algae cultures ( approximately 1x104 cells per flask). Following 72-hours incubation, algal cell densities were determined using a hemacytometer. Values were 127,76,109,69 and 1%, respectively, of the solvent control response. These values were used to set exposures for the definitive phase.

In the definitive phase, triplicate algal cultures were exposed to the test substance at nominal concentrations of 16, 31, 63,130, 250, and 500 ug/L (ppb). Cell densities were monitored at 24-, 48-, and 72-hours following study initiation. From these data, EC50 (50% decrease) values for Biomass (EbC50) and Growth Rate (ErC50) were calculated. Test substance concentrations in the test media were determined at 0- and 72-hours using HLPC. The mean concentrations were 7.5, 13, 14, 28, 50, and 79 ug/L (ppb). The inhibitions of algae Growth Rates for the test substance in the definitive 72-hour study were 0, 2, 15, 20, 32, and 38% (relative to pooled control values) for the measured test substance concentrations of 7.5, 13, 14, 28, 50, and 79 ug/L (ppb). Corresponding inhibitions of Biomass generation were 15, 41, 59, 63, 81, and 91%. Individual cell appearances were found microscopically to be normal for surviving cells except cellular bloating was noted at the highest exposure level. Calculations indicated that the ErC50 for the test substance was > 79 ug/L (ppb) while the EbC50 was 18 ug/L (ppb). The No Observed Effect Concentrations (NOECs) were assumed to be equivalent to EC10 values, and accordingly were EbC10 = 4.3 ug/L (ppb) and

The EC50 values for the test substance ranged from 18 to > 79 ug/l (ppb) for Biomass increases and Growth Rates. The NOECs ranged from 4.3 to 31 ug/L (ppb) for these parameters.

Test substance: Tested as the commercial product Reliability: (1) valid without restriction

ErC10=31 ug/L (ppb).

31-JUL-2000 (30)

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Date: 30-OCT-2001 ID: 68953-84-4 5. Toxicity

#### 5.1 Acute Toxicity

5.1.1 Acute Oral Toxicity

Type: T<sub>1</sub>D50 Species: rat Strain:

Result:

Sex: no data

Number of
Animals:
Vehicle:

Value: > 2000 mg/kg bw

Method: other: Directive 84/49/EEC, B.1

Year: 1990 GLP: yes

Test substance: as prescribed by 1.1 - 1.4
Reliability: (1) valid without restriction

01-AUG-2000 (7)

Type: LD50 Species: rat

Strain:

Sex: male/female

Number of

Animals: 10

Vehicle: other: corn oil Value: > 5000 mg/kg bw

Method: other: US EPA 40CFR798.2650, Oral Toxicity-Limit Test

Year: 1993 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: Five (5) male and five (5) female young adult rats

(Sprague-Dawley) were administered a single dose of the test substance by gavage. The test substance was dispersed in corn oil (Sigma Chemical Company) and administered at a dosage of 5000 mg/kg. The animals were observed for clinical signs of toxicity at approximately 1-, 4- and 24-hours following administrations on the day of dosing and daily thereafter for 14-days. Body weights were recorded on Day-0, Day-7 and Day-14. All animals were subjected to a gross

necropsy at study termination.

Result: One (1) animal died during the 14-day observation period.

Clinical signs observed included decreased activity, decreased muscle tone, and diarrhea. No significant impairment on body weight gains were noted in either the male or female rats. Necropsy of the animal that died during the study revealed discolored kidneys, spleen, and liver. No visible lesions were observed in any of the animals at terminal necropsy. The estimated acute oral LD50 (combined

sexes) for the test substance was determined to be > 5000

mg/kg.

Reliability: (1) valid without restriction

01-AUG-2000 (20)

5.1.2 Acute Inhalation Toxicity

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Date: 30-OCT-2001 ID: 68953-84-4

5. Toxicity

Type: LD50 Species: rabbit

Strain:

Sex: male/female

Number of

Animals: 10 Vehicle: other

Value: > 2000 mg/kg bw

Method: OECD Guide-line 402 "Acute dermal Toxicity"

Year: 1995 GLP: yes

Test substance: other TS

Method: Albino rabbits (five males and five females) were shaved in

the caudal portion of the animals' trunks. One (1) day

later, a 2000 mg/kg dose of 40 mesh test substance (obtained by grinding in motar/pestle) was placed onto the skin sites

(approximately 10% of the body surface areas). The

application sites were then covered with gauze, plastic, and elastic wraps and finally secured with non-irritating tape. After 24-hours of skin contact to the exposure areas, the gauze patches were removed and adhering test substance removed with moistened gauze. Skin test sites were scored for signs of erythema (redness) and edema (swelling) according to Draize procedures from Day-1 to Day-14

following cessation of exposures. Animals were observed for adverse clinical signs, mortality, and body weights (Day-0, Day-7, and Day-14). Necropsies were performed on the final

day of observations (Day-14).

Remark: A limit test

Result: The test substance induced no deaths or apparent adverse

clinical signs. Mild irritation (Grades 1,2 erythema; Grade 1 edema) was seen at skin sites of treated rabbits for periods ranging fron Day-1 to Day-10. Staining of skin was noted due to the dark color of the test substance. A body weight decrease was seen in one (1) of the ten (10) rabbits between Day-7 and Day-14. No compound-related non-dermal findings were observed in the study. No mortality or adverse clinical/necropsy changes were observed associated with the test substance. The dermal LD50 for the test substance was

shown to be > 2000 mg/kg.

Test substance: Tested as the commercial product Reliability: (1) valid without restriction

01-AUG-2000 (26)

#### 5.1.4 Acute Toxicity, other Routes

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Date: 30-OCT-2001 ID: 68953-84-4

#### 5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: Fischer 344
Route of admin.: oral feed
Exposure period: 28 days

Frequency of

treatment: Daily

Post. obs.

period: 2 weeks

Doses: 0, 7.5, 30 and 120 mg/kg/day

Control Group: yes, concurrent vehicle

NOAEL: 7.5 mg/kg LOAEL: 30 mg/kg

Method: other: Oral 4-week dietary study
Year: 1996 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: The tes

The test substance was prepared by grinding in a coffee mill, sieved through a 125 um mesh screen and mixed with rodent diet NIH-07 at 0, 120, 470, 1900 ppm (0, 7.5, 30, and 120 mg/kg/day). Stability, homogenity, and dose verification were performed to confirm compliance with protocol. The prepared dosed feed was presented to 14 male and 14 female rats (Fischer 344) per test group at twelve weeks of age for four (4) weeks. Six (6) rats/sex/group were held for post-exposure in two (2) week recovery groups. Test rats were monitored for body weights, feed consumption, and clinical signs. Collections were performed on six (6) or three (3) rats/sex/group at 28-days and 42-days sacrifice periods for blood (hematologies and clincal chemistries) and urinalyses, respectively. Necropsies were performed on all rats, and organs were weighed (liver, kidneys, pituitary, uteri, heart, brain, spleen, thryoids, adrenals, testes, and ovaries). These and other major organs were preserved in formalin, stained with H&E, and subjected to microscopic evaluations. Liver, kidney, and urinary bladder slices were subjected to immunohistochemical staining for proliferating cell nuclear antigen (PCNA) for assessment of cellular division.

Result:

The test substance was shown to be completely stable in diets for 46-days. Mixing procedures produced homogeneous diets that were found within 10% of target concentrations. No compound-related deaths occurred, The boby weights were not affected in male rats whereas the high dose female rats displayed 5% body weight decreases during study weeks two (2) through four (4). Food consumption was decreased in the high dose males and in the mid- and high dose females mainly during study weeks two (2) through four (4). Various test substance-induced hematological changes occurred that included: increased mean corpuscular volumes and decreased mean corpuscular hemoglobin concentrations (high dose males and females) and blood bilirubin and cholesterol increases (high dose males and females). Most blood endpoints tended to approach control levels during week two (2) of the recovery

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weight increases were seen at 28-days for liver and kidneys (high dose males and females; mid-dose females) and heart and spleen (high dose females). Only the kidney weights did not reach control levels by 42-days. There were no gross tissue or microscopic changes related to the test substance. Proliferating cell nuclear antigen (PCNA) exams showed cell division changes for: increases for liver cells (High dose males and females and mid-dose males at 28-days only); changes for kidney cells (decreases in high dose females at 28-days and increases in high dose males and females at 42-days; and increasing trend in urothelial cells in bladder (low and mid-dose males and females at 28-days). Macrocytic anemia was the primary change in rats related to the test substance administration. This change was reversible within 2 weeks following dietary exposure as were liver weight and serum cholesterol elevations. These changes were very minor, and had no apparent toxicological significance in this study. The lack of dose-responsiveness in the PCNA data provides results of uncertain importance to the assessment of the toxicity of this test substance.

Reliability: (1) valid without restriction

02-AUG-2000 (11)

Species: rat Sex: male/female

Strain: other: Fischer 344/N TacfBR

Route of admin.: gavage Exposure period: 21 days

Frequency of

treatment: Daily

Post. obs. period:

Doses: 0, 0.1, 0.3, 1.0, and 3.0 g/kg/bw

Control Group: yes, concurrent vehicle

LOAEL: 100 mg/kg bw

Method: other: Oral 3-Week Range-Finding Study Year: 1994 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: A 4-week diet-study was also conducted.

Result: Doses of 1.0 and 3.0 g/kg/day of WINGSTAY 100 administered

by gavage for up to 6 days were lethal for male and female

F344 rats. The only pertinent gross finding of all unscheduled deaths was the paleness of most external surfaces and viscera. The mid-low (0.3~g/kg/day) and low

(0.1 g/kg/day) doses caused time and dose related

significant body weight gain loss, liver weight increase and

hepatocellular labeling index increase at 0.1 g/kg.

Therefore, in the subchronic studies, the recommended daily dose of WINGSTAY 100 should not exceed 100 mg/kg/day, if

administered by gavage.

Test substance: The test substance was prepared in an olive oil suspension

for dosing

Reliability: (1) valid without restriction

02-AUG-2000 (5)

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Date: 30-OCT-2001
5. Toxicity ID: 68953-84-4

#### 5.5 Genetic Toxicity 'in Vitro'

Type: Ames test

System of

testing: Ames/E. coli preincubation; Salmonella typhimurium TA-98, 100,

1535, 1537, 1538, and WP2 uvrA

Concentration: Salmonell stains without S9 activation: 0.167, 0.5, 1.67, 5,

16.7, and 50 ug/plate; Salmonella strains with S9

activation:1.67, 5, 16.7, 50, 167, and 500 ug/plate; E.coli with/without S9 activation:1.67, 5, 16.7, 50, 167, and 500

ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: positive

Method: other: Japan's Industrial Safety & Health Law, a combination

of OECD Guidelines 471 and 472.

Year: 1993 GLP: yes

Test substance:

Method:

as prescribed by 1.1 - 1.4
In a preliminary assay, revertant frequencies for all doses

of the test substance in tester strains TA1535, TA1537, TA98, TA100, and WP2 uvrA with S9 metabolic activation, and in tester strains TA1535, TA1538, TA98, TA100 and WP2 uvrA without S9 activation, approximated the concurrent negative controls. However, statistically significant, increases in revertant frequencies, to approximately 1.7- to 2.5-fold control values, were observed in tester strain TA1538 with S9 metabolic activation and in tester strain TA1537 without

S9 metabolic activiation. In addition, the increases

observed in strain TA1538 with S9 metabolic activation were

dose dependent.

In a confirmatory assay, revertant frequencies for all doses of the test substance in tester strains TA1535, TA100, and WP2 uvrA with metabolic activation, and in tester strains TA1535, TA1538, TA98, TA100, and WP2 uvrA without S9 metabolic activation, approximated control values. Statistically significant, dose-dependent increases in revertant frequencies, to control values, were observed in tester strains TA1537, TA1538, and TA98 with metabolic activation. Statistically significant increases in revertant frequencies, to control values, also were observed in tester strain TA98 without S9 metabolic activation. However, these latter increases apparently were not dose related.

The test substance was re-evaluated in all five Salmonella strains with S9 metabolic activation, and in tester strain TA98 without S9 metabolic activation. Revertant frequencies for all doses of the test substance in tester strains TA1535, TA1537, and TA100 with S9 metabolic activation, and in tester strain TA98 without S9 metabolic activation, approximated or were less than control values. Statistically significant, dose-dependent increases in revertant frequencies, to control values, were observed in tester strains TA1538 and TA98 with S9 metabolic activation. All

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within acceptable limits.

Result: The test substance was shown to cause mutations in

Ames/Salmonella strains TA1538 and TA98 with S9 activation.

positive and negative control values in all assays were

(1) valid without restriction Reliability:

04-AUG-2000 (16)

Type: Ames test

System of

testing: Ames/Salmonella-E.coli Liquid Pre-incubation Assay in

Salmonella strains TA1535, TA1537, TA1538, TA98, and TA100 and

in E.coli strain WP2 uvrA.

Salmonella strains with S9: 1.67, 5, 16.7, 50, 167, and 500 Concentration:

ug/plate; Salmonella strains without S9: 0.167, 0.5, 1.67, 5, 16.7, and 50 ug/plate; E.coli with/without S9: 1.67, 5, 16.7,

50, 167, and 500 ug/ plate.

Cytotoxic Conc.:

Metabolic

with and without activation:

positive Result:

Method: other: Japan's Industrial Safety & Health Law, a combination

of OECD Guidelines 471 and 472.

1994 GLP: yes

Test substance:

as prescribed by 1.1 - 1.4

In a preliminary assay, revertant frequencies for all doses Method:

of the test substance in tester strains TA1535, TA1537, TA100, and WP2 uvrA with and without S9 metabolic activation approximated the concurrent negative controls. However,

statistically significant, increases in revertant

frequencies, to control values, were observed in tester strains TA1538 and TA98 with S9 metabolic activation. In addition, the increases observed in strain TA1538 with S9

metabolic activation were dose dependent.

In a confirmatory assay, revertant frequencies for all doses of the test substance in tester strains TA1535, TA100, and WP2 uvrA with metabolic activation, and in tester strains TA1535, TA1537, TA1538, TA100, and WP2 uvrA without S9 metabolic activation, approximated control values.

Statistically significant, dose-dependent increases in revertant frequencies, to control values, were observed in tester strains TA1537, TA1538, and TA98 with metabolic activation. Statistically significant increases in revertant frequencies, to control values, also were observed in tester strain TA98 without S9 metabolic activation. However, these latter increases apparently were not dose related.

The test substance was re-evaluated in all five Salmonella strains with S9 metabolic activation, and in tester strain TA98 without S9 metabolic activation. Revertant frequencies for all doses of the test substance in tester strains TA1535, and TA100 with S9 metabolic activation, and in tester strain TA98 without S9 metabolic activation, approximated control values. Statistically significant, dose-dependent increases in revertant frequencies, to

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control values, were observed in tester strains TA1537, TA1538, and TA98 with S9 metabolic activation. All positive and negative control values in all assays were within

acceptable limits.

Result: The test substance was shown to cause mutations in Ames/Salmonella strains TA1537, TA1538 and TA98 with S9

metabolic activation.

Reliability: (1) valid without restriction

04-AUG-2000 (17)

Type: Cytogenetic assay

System of

testing: Chromosomal aberration assay in CHO cells

Concentration: 0.4, 2, 4, and 25 ug/mL

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: OECD Guide-line 473 "Genetic Toxicology: In vitro Mammalian

Cytogenetic Test"

Year: 1993 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: In the structural Chromosomal Aberration assay, duplicate cultures were established for each dose level. Three

treatment schedules were used: a) First set of cultures were treated for 5-hours with the appropriate dose of the test sample in Ham's F12 serum free (F12SF) medium either in the presence or absence of S9 metabolic activation along with concurrent negative and positive controls followed by three (3) Puck's saline washes and medium replacement; b) Second set of cultures were treated for 24-hours with the test

substance or control articles in Ham's F12 medium containing five (5) % serum (F12FCM5%) without S9 metabolic activation, and; c) Third set of cultures were treated for 48-hours with

the test substance or control articles in F12FCM5% medium without S9 metabolic activation. Two (2) to three (3) hours prior to harvest, Colcemid (2X10-7M) was added to all sets of cell cultures to arrest dividing cells in metaphase. CHO cells were harvested at the eppropriate time and metaphase slides were prepared and stained.

The data from one hundred metaphases from each culture (200 metaphases per dose point) were pooled for statistical analysis. Data were evaluated by using the chi-square of aberrant versus normal cells while comparing each dose level to its concurrent negative control. The data were also analyzed for statistical significance by pairwise t-tests comparing the number of aberrations per cell in each treated dose versus the negative control.

Analysis of the data for the 24-hour treatment with the test substance indicated that there were statistically significant dose-related increases in the frequency of aberrations/cell and proportion of aberrant metaphases at doses 2 and 4 ug/mL. The data for the 2 and 4 ug/mL doses produced a statistically significant linear trend when

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Result:

5. Toxicity

analyzed by the Cochran/Armitage Linear Trend Test. To verify the biological sighificance of this finding, the 24-hour treatment was repeated.

In the confirmatory assay, the test substance was re-evaluated at doses of 25 ug/mL with S9 metabloic activation (5-hour treatment) and 0.4, 2, and 4 ug/mL without S9 metabolic activation (24-hour treatment). Analysis of the data for the 5-hour treatment did not produce statistically significant increases in aberrations/cell or in proportion of aberrant metaphases.

Analysis of the data for the 24-hour treatment indicated a statistically significant increase in aberrations/metaphase at the mid-dose (2 ug/mL) with S9 metabolic activation but there were no significant increases in the proportion of aberrant metaphases. However, when the data for 2  $\mbox{ug/mL}$ (0.045 + or - 0.208) were compared to the untreated control data (0.025 + or - 0.157) or to Pharmakon historical acetone data (0.034 + or - 0.021), there were no statistically significant increases in the frequency of aberrations/metaphase. Therefore, the positive finding in the t-test for 2 ug/mL was considered a statistically artifact with no biological significance. There were no other statistically significant increases in aberration/metaphase or in the proportion of aberrant metaphases at any of the remaining dose levels for the 24-hour treatment.

The test substance was judged negative (non-clastogenic) based on its inability to reproducibly induce dose-related increases in structural chromosomal aberrations in CHO

cells.

Reliability: (1) valid without restriction

20-FEB-2001 (19)

Type: DNA damage and repair assay

System of

testing: E. coli Pol A1- Liquid Suspension Assay

Concentration:
Cytotoxic Conc.:

Metabolic

activation: without
Result: positive
Method: other

Year: 1980 GLP: no

Test substance: as prescribed by 1.1 - 1.4 Reliability: (2) valid with restrictions

Although the study was old and was not conducted to GLP, the

test parameters were based on a scientifically sound procedure for that time period and the study was properly

conducted.

04-AUG-2000 (32)

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Type: other: Transformation Assay

System of

testing: Balb/3T3 In Vitro Transformation Assay

Concentration: .01 ug/ml to 1.0 ug/ml

Cytotoxic Conc.:

Metabolic

activation: without
Result: negative
Method: other

Year: 1981 GLP: no

Test substance: as prescribed by 1.1 - 1.4 Reliability: (2) valid with restrictions

Although this study was probably not conducted to GLP, the

test parameters used were based on a known and well

established procedure.

04-AUG-2000 (12)

Type: other: Unscheduled DNA Synthesis Assays (UDS) with Rat

Hepatocytes

System of

testing: Hepatocytes form male Fischer 344 (F344/Crl) rats

Concentration: Slightly above their limits of solubility

Cytotoxic Conc.:

Metabolic

activation: without Result: negative

Method: other: Unscheduled DNA Synthesis Assays (UDS) with Rat

Hepatocytes on Test substance Condensation Products

Year: 1999 GLP: yes

Test substance: other TS: Test substance condensation products with

Dicylopentadiene

Method: The test substance, 1,4-Benzenediamine, N.N'-mixed Ph and

tolyl. derivs., was reacted with Dicyclopentadiene in varying ratios, resulting in three condensation products. Each of these condensation products were subjected to independent in vitro unscheduled DNA synthesis (UDS) assays with hepatocytes form male Fischer 344 (F344/Crl) rats. All

three (3) condensation products were tested at

concentrations slightly above their limits of solubility in the tissue culture medium. Hepatocytes were exposed to test substances for 18-20 hours to allow bioactivation and DNA

repair. The assay was based on the incorporation of 3H-thymidine into the hepatocyte's DNA during repair of DNA-damage. This incorporation was monitored by counting Net

Nuclear Grains (NNG) formed on photographic emulsion placed on the cells adhering to glass slides. Criteria for a positive response included: (a) Significant increase in number of grains at two (2) levels of exposure above

negative control levels, (b) A dose-responsiveness in grain counts up to toxic levels of exposure, and (c) At least one (1) value for NNG that is five (5) or above. A negative

response is reported for NNG's that are <0, and an equivocal

or inconclusive response are results that are 0<#<5.

Result: In all the Unscheduled DNA Synthesis Assay (UDS) trials, the

three (3) negative controls {the untreated cells control, F,

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and Dimethylsulfoxide (DMSO)} had negative values for Net

Nuclear Gain (NNG) counts (<0). A positive control, 2-Aminofluorene (2-AF) was positive for induction of UDS; the mean NNG counts were 45.92 and 58.99 in the first and second assays, respectively, indicating assay validity. (i.e., hepatocytes were capable of metabolic activation and DNA repair). The positive control responses occurred at toxic levels. UDS assay results for NNGs were in the range of -26 to -46, demonstrating a lack of UDS activity for the three (3) condensation products at concentrations greater than their solubilites in the test media. The results indicated that, under controlled laboratory conditions, the condensation products from the reaction of 1.4-Benzenediamine, N,N', mixed Ph and tolyl. derivs. with Dicyclopentadiene were negative for induction of UDS in rat hepatocytes at concentrations up to and greater than their solubilities. This assay demonstrated a lack of genetic activity in this mammalian DNA-repair test system.

Reliability: (1) valid without restriction

07-AUG-2000 (36)

#### 5.6 Genetic Toxicity 'in Vivo'

Type: Drosophila SLRL test

Species: Drosophila melanogaster Sex:

Strain:

Route of admin.: oral feed Exposure period: 24 hours

Doses: 50 ug/ml and 10 ug/ml

Result: negative

Method: other: Drosophila melanogaster (Fruit Fly) System

Year: 1979 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Result: Negative under conditions of the assay

Reliability: (2) valid with restrictions

Although the study was old and was not conducted to GLP, the

test parameters were based on a scientifically sound procedure for that time period and the study was properly

conducted.

04-AUG-2000 (31)

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Type: Drosophila SLRL test

Drosophila melanogaster

Species: Strain:

Route of admin.: oral feed Exposure period: 24 hours

Doses: 0.05 mg/ml and 0.63 mg/ml

Result: negative

Method: other: Drosophilia SLRL Assay

Year: 1979 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Result: Negative under conditions of the assay.

Reliability: (2) valid with restrictions

Although the study was old and was not conducted to GLP, the

Sex:

test parameters were based on a scientifically sound

procedure for that time period and the study was properly conducted.

04-AUG-2000 (13)

Type: Micronucleus assay

Species: mouse Sex: male/female

Strain: CD-1 Route of admin.: i.p.

Exposure period: single dosing

Doses: 0, 250, 1250, 2500 mg/kg test chemial; 0.5 g/kg TEM (+

control)

Result: negative

Method: OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test"

Year: 1993 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: Nine (9) groups of mice (CD-1) were acclimated to laboratory

conditions for 25-days prior to initiation of the study. THe mice were randomized by body weight and assigned to groups

using a computer-generated random number list.

Each group of mice was comprised of ten (10) animals(five (5) males/five (5) females). Each mouse received a single intrperitoneal dose at 10 mL/kg of body weight. The test substance at dose levels of 250, 1250, anf 2500 mg/kg was

adnimistered to three (3) groups of mice which were sacrificed at 24-, 48-, and 72-hours post dose.

Concurrently, the negative control, Dimethylsulfoxide (DMSO)/corn oil, was administered, as dose volume of 10 mL/kg of body weight, to three (3) groups of mice. A group of these mice were included in each sampling time. The positive control, Triethylenemelamine at 0.5 mg/kg, was administered to one (1) group of mice and sacrificed at

24-hours post dose.

All mice were sacrificed and their femurs were removed. Their bone marrow was removed by flushing. Smears were made of the suspended cells.

One (1) thousand young erythrocytes were evaluated for a change of ratio of polychromatic erythrocytes (PCE) to normochromatic cells (NCE).

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5. Toxicity

Result:

There were no statistically significant depressions in the PCE/NCE ratios in any groups of mice except for the 2500 mg/kg group at 48-hours sacrifice time (p<0.01) which was an indication that the test substance had reached the bone marrow and was toxic to erythrocytes.

Analysis of the micronucleus data for the groups treated with the test substance indicated that there were no statistically significant increases in the frequency of

micronucleated PCEs. The test substance was judged negative

(non-clastogenic) based on its inability to induce

micronucleated PCEs.

Reliability: (1) valid without restriction

04-AUG-2000 (18)

Type: other: 32P Postlabeling Assay for Detection of Adduct

Formation in Rat DNA

Species: rat Sex: male/female

Strain: other: Fischer 344/N TacfBR

Route of admin.: gavage Exposure period: 7 days

Doses: 0., 0.3, 1.0, and 3.0 g/kg/bw

Result: negative

Method: other: 32P Post-Labeling Assay for DNA Adduct Formtion

Year: 1995 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Remark: The purpose of the study was to determine the potential of

WINGSTAY 100 to bind covalently to liver and urinary bladder DNA of male and female rats after in vivo administration of

WINGSTAY 100.

Result: Under conditions of the study, the test substance did not

induce DNA-adducts in the liver and urinary bladder DNA of

rats.

Reliability: (1) valid without restriction

07-AUG-2000 (4)

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Date: 30-OCT-2001
5. Toxicity ID: 68953-84-4

#### 5.8 Toxicity to Reproduction

Type: Two generation study

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: oral feed

Exposure Period: F0 exposed during 10 weeks premating, 2 weeks of mating, 3

weeks (gestation), and through the weaning (21 day) period. F1 males and females exposed for 10 weeks prior to mating.

Frequency of

treatment: Daily
Premating Exposure Period
male: 10 weeks
female: 10 weeks
Duration of test: 9 months

Doses: 0, 120, 400 or 1500 ppm. Control Group: yes, concurrent no treatment

Method: OECD Guide-line 416 "Two-generation Reproduction Toxicity

Study"

Year: 2000 GLP: yes

Test substance: as prescribed by 1.1 - 1.4

Method: This study was designed in compliance with EPA GLP and USEPA

FIFRA guidelines. Dose levels were established from a rangefinding study at Research Triangle Institute which employed dietary levels of 120, 1900, and 5700 ppm of WINGSTAY 100. The top level was lethal to dams and

offspring, 1900 ppm induced one nonviable litter in 9 total,

and thus, the top dose for the definitive study was

decreased by 20% to assure high viability in test group. No

effects were seen at 120 ppm.

This study used 30 SpragueDawley rats/sex/dose (F0) exposed to diets containing 0, 120, 400 or 1500 ppm WINGSTAY 100 during 10 weeks premating, 2 weeks mating, 3 weeks

(gestation), and through the weaning (21 day) period. F1 litters were culled to 10 each each at 4 days postnatal (PND) 30 other F1 males and females/group chosen for

pairing, and fed WINGSTAY 100 as above for 10 weeks prior to mating. After mating/gestation of F1, the resulting F2 rats were delivered, and maintained through weaning period (to PND 21). Weekly body weights (BWs) and food consumption (FC), and daily clinical observations were recorded. Necropsies and histopathology (primary kidneys) were performed on selected rats from each sex/group/generation (all F0 and F1 dams at PND21, three F1 and F2 pups/test

group at PND21). Remaining F1 and F2 rats were euthanized without examination. Data were collected on vaginal cytology, mating, pregnancy, litter, and pup parameters.

WINGSTAY 100 induced dystocia (difficult deliveries) in pregnant rats which may have led to prolonged gestation and

increased perinatal deaths, decreased live births, and increased pup weights. In addition, polycystic lesions were

observed at all dose levels. Prolonged gestation has

previously been associated with the WINGSTAY component DPPD,

and polycystic kidneys were observed in DPamine-treated

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Remark:

rats. Based upon adult toxicities, reproductive and offspring endpoints, there was no NOEL for WINGSTAY 100 in

this study.

Result: High dose females had decreased Body Weights (BWs) relative

to other test groups throughout majority of study period. Mortality during gestation/lacation were: F0 dams- 0 in 24 pregnancies, 0/27, 3/24, 4/25; F1- 0/22, 0/23, 1/22, 1/24. Numbers of pregnancies with no live births: F0-0, 1, 1, 10; F1- 0, 1, 1, 2. Gestational length: F0- 22.2 days, 22.4 days, 22.8\*, 23.5\*; F1- 22.2, 22.8\*, 23.1\*, 23.2\* (\* = statistically sighnificant). The number of live pups/litter: F0-15.6, 14.1, 11.9, 7.6\*; F1- 15.6, 13.7, 13.3, 10.8\*. Pups weights (g) on PND 0: F0-6.38, 6.79\*, 6.93\*, 6.63\*; F1-6.32, 6.89\*, 6.99\*, 6.63\*. WINGSTAY 100-related kidney lesions were observed grossly (as white or clear cysts) and microscopically (polycyctic findings with variable severity): F0 adults- males 0/0, 0/0, 0/0, 0/1 and females 0/0, 0/0, 0/2, 3/9; F1 weanlings- males 0/23, 1/25, 8/20, 10/11 and females 0/22, 5/26, 7/18, 11/11; F1 adults- males 0/30, 5/30, 10/30, 21/30 and females 0/30, 2/30, 1/30, 18/30; F2 weanlings- males 0/60, 3/64, 6/19, 15/16 and females 0/60, 5/64, 8/19, 15/15. The severity of kidney

lesions were also dose related.

Reliability: (1) valid without restriction

11-FEB-2001 (35)

# 5.9 Developmental Toxicity/Teratogenicity

Species: rat Sex: female

Strain: Sprague-Dawley

Route of admin.: gavage Exposure period: 10 days

Frequency of

treatment: Dosed on days 6-15 gestation

Duration of test:

Doses: 0, 20, 70, 200 mg test material in 5 ml corn oil/kg

Control Group: yes, concurrent vehicle

NOAEL Maternalt.: 70 mg/kg bw NOAEL Teratogen.: <= 200 mg/kg bw

Method: OECD Guide-line 414 "Teratogenicity"
Year: 1995 GLP: yes

Test substance: other TS

Method: Preliminary trials in 8 rats/group indicated that 600 mg/kg

was lethal to 50% of maternal rats while 200 mg.kg caused decreased body weights in materanl and fetal animals. There were no effects at 20 or 70 mg/kg. Consequently, 200 mg/kg was selected as the top (high) dose in the definitive study,

Confirmation of the test dose solutions were confirmed

analytically.

The definitive study used 25 inseminated female rats per test group (0, 20, 70, and 200 mg of test substance/kg doses in five (5) mL corn oil/kg). The animals were dosed on Days 6-15 gestation. Body weights, food consumption, liver

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weights, clinical changes, pregnancy rates, and corpora lutea counts were followed along with numerous fetal parameters. All fetuses were weighed, sexed, and assessed for external and visceral abnormalities. One (1) half of the fetuses were examined for skeletal abnormalities while the second half were subjected to cranial bone assessments.

Remark: Administered in 5 ml corn oil/kg by gavage

Result: The test substance induced no lethality. Deficits were seen

in maternal body weights (Day-12 and body weight change from Day-6 to Day-15) and food consumption (during treatment period) at the highest dose only (200 mg/kg). Pregancy rates, litter sizes, number of live fetuses, uterine

implantation, and all gestational parameters were unaffected by chemical treatment. There was a linear trend towards lower body weights in fetuses with increasing doses

(approximately 5% decrease in 200 mg/kg group). Assessment of cranial, skeletal, visceral, and external appearance discerned no compound-related abnormalities (malformatiuons or variations) according to established criteria. The test material produced minimal effects (body weight) to maternal rats from oral dosing of 200 mg/kg during pregnancy. There was no induction by the test chemical of birth defects

(major or minor) in fetal animals. Tested as the commercial product (1) valid without restriction

08-AUG-2000 (21)

Species: Sex: male/female rat

Strain: Sprague-Dawley Route of admin.: oral feed

Exposure period: Varied, see method

Frequency of

Test substance:

Reliability:

treatment: Varied, see method

Duration of test:

2500 ppm Doses:

Control Group: yes, concurrent vehicle Method: other: Mechanistic Study

Year: 2000 GLP: no

Test substance: as prescribed by 1.1 - 1.4

Method: The toxicity of the test substance to maternal and 1st.

generation offspring was evaluated by exposing CD

(Sprague-Dawley) rats to fixed dietary concentrations of 2500 ppm during different time periods (i.e. exposures during prebreed, mating, gestation, and/or lactation). Five (5) Groups (20/sex/Group) were studied including: Group one (1) - Negative control; Group two (2) - Dietary test substance during prebreed and mating, exposures ended on gestation day

(qd)-0; Group three (3)- Dietary test substance during gestation and lactation, exposures began on qd-0; Group four

(4) - Dietary test substance during prebreed, mating, gestation, and lactation, the Positive control and; Group five (5)- Dietary test substance during prebreed, mating, gestation, and lactation, plus 600 ppm of iron gluconate in the drinking water for prebreed through lactation.

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Males and females were paired within Groups (1:1) for the two-week mating period. Once a given female was found to be sperm positive {date designated as gestation day (gd)-0}, "her" male was euthanized and discarded. On the day of delivery (pnd-0), pups were counted, sexed, and weighed. On pnd-4, litters were culled to ten, counted, sexed, and weighed. On pnd-7, -14, and -21, pups were counted, sexed, and weighed. All pups were euthanized and one (1)/sex/litter necropsied on pnd-21. Dead pups on pnd-0 and -1 were examined macroscopically (necropsied) for polycystic kidneys. Female body weights and feed consumption were recorded weekly during prebreed, gestation, and postnatally. At necropsy on pnd-21, the maternal spleen, liver, and kidneys were weighed and retained in a fixative. Kidneys form Groups one (1) and five (5) were examined histopathologically. Blood sampling was performed on gestation day-21 and pnd-21 from all females (pregnant) by tail vein withdrawal. Blood sampling was performed on pnd-21 on the F1 offspring by withdrawal from the abdominal vena cava at sacrifice. The blood parameters assessed were: WBC, RBC, Hgb, Hct, MCV, MCH, MCHC, RDW, Platelets, WBC Diffential ( to correct the RBC and WBC counts for Nucleated Red Blood Cells) and Methemoglobin. On gd-21, a second sample of blood was taken via tail vein from all pregnant females in all Groups, with plasma frozen for possible subsequent analysis for specific hormones. For Group three (3), any female who had not yet delivered by gestation day-23 had blood taken from the tail vein and plasma frozen. On pnd-21, the spleen, liver, kidneys, and heart from one (1) pup/sex/litter were weighed and retained in a fixative. The kidneys from all offspring were examined histologically. Statistical analysis included both parametric and nonparametric tests for continuous and discrete data. The objectives of this study were to confirm and further characterize prviously-observed effects following the test substance administration to pregnant rats. This study was designed (1) to determine the necessary and sufficient timing of exposure to maternal females at a fixed dietary concentration of the test substance to produce dystocia, prolonged gestation, and polycystic kidneys in offspring, (2) to determine whether the test substance results in demonstratable macrocytic anemia in maternal animals, (3) to determine if there is treatment-induced anemia and whether iron supplementation ameliorates or prevents the anemia, dystocia, and/or polycystic kidneys, and (4) to determine if FO parental females exhibit polycystic kidneys due to

Remark:

Result:

dietary exposure to the test substance. FO Males: The test substance intake over the prebreed period (Study Days 0-28) averaged 180 mg/kg/day for all three (3) exposed Groups {two (2), four (4), and five (5)}. Iron gluconate intake in Group five (5) averaged 56 mg/kg/day (Study Days-0 to 28). Clinical observations were found to be unrelated to compound administration.

F0 Females: The test substance intake averaged 187-192

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> mg/kg/day for Groups two (2), four (4) and five (5) during gestation days (gd)-0 to 28. Iron gluconate intake during gestational days-0 to 28 in Group five (5) averaged 53 mg/kg/day. Clinical observations during gestation included one (1) female found dead in Groups three (3) and four (4), alopecia predominantly in Groups four (4) and five (5), pale eyes and tail, pale (not otherwise specified) almost exclusively in Groups three(3), four (4) and five (5) (all exposed), pilorection in Groups three (3), four (4) and five (5), and delayed parturition in Groups three (3), four (4), and five (5). The hematological profile of maternal rats on gestation day-21 found no evidence on macrocytic anemia in any groups.

> REPRODUCTIVE/DEVELOPMENTAL: Gestational index (a measure of live litters relative to pregnant females) was significantly increased in Groups three (3) and four (4) but not in Group five (5). Male mating, fertility, and pregnancy indices were equivalent across all groups. Gestational length in days was significantly prolonged in Group three (3) (23.6+/-0.2), Group four (4) (23.8+/-0.2), and Group five (5) (23.5+/-0.2)relative to Control Group value (22.2+/-0.1) and the value in Group two (2) (22.3+/-0.1). Number of implantation sites per litter was significantly reduced in Group five (5). Percent of postimplantation loss was significantly increased in Groups three (3) and four (4). Pups per litter were significantly reduced in Groups three (3), four (4) and five (5), and number of dead pups per litter were significantly increased in Groups three (3) and four (4). Weanling gross and microscopic findings were limited to hydronephrosis in Groups one (1) and two (2), gas in intestines in Group two (2), and gross evidence of polycystic kidneys in Groups three (3), four (4), and five (5). Maternal hematologic profiles at sacrifice (21 days after delivery) indicated statistically significant changes in most erythrocyte parameters. The white blood cell differential counts indicated changes (as percent of cells examined) as follows: increase in segmented neutrophils and decrease in lymphocytes only in Group four (4), with no treatment-related changes in the percentages of monocytes or eosinophils. Histopathologic assessment was performed on

kidneys of all maternal rats in Groups one (1) and five (5). Polycystic kidneys were observed microscopically (but not macroscopically) in three (3) of 20 animals in Group five (5), with no polycystic kidneys observed in Group one (1).

The timing of exposure to the test substance with respect to pregnancy is an important determinant of toxicity. Exposure of F0 females to 2500 ppm of the test material during gestation is necessary and sufficient to produce dystocia (prolonged gestation). It is necessary and sufficient to expose F0 dams during gestation and/or lactation to produce polycystic kidneys in the F1 offspring. Since no Groups were exposed only during gestation or only during lactation, it is not possible to further define how exposure timing

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#### 5. Toxicity

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affects this endpoint. There was no demonstratable macrocytic anemia in gestation day-21 (gd-21) F0 dams in any treatment Group, but at post delivery day-21 (pnd-21), F0 mothers exposed prior to and during mating, gestation, and lactation were anemic. The F1 offspring at pnd-21 did not consistently display evidence of macrocytic anemia. Iron supplementation did not affect pnd-21 maternal anemia, dystocia, or incidence/severity of polycystic kidneys in the F1 offspring. However, perinatal survival of the offspring was affected. Microscopic, but not macroscopic evidence of polycystic kidneys was found in 15 percent of dams treated prior to and during mating, gestation, and lactation (with iron supplementation). Controls had neither macroscopic nor microscopic indications of polycystic kidneys. Exposure of animals to the test substance prior to and during mating {Group two (2)} did not appear to result in adverse affects to offspring. Furthermore, exposure during the prebreed/mating periods did not increase the affects produced from gestation/lactation exposures only.

Reliability:

(2) valid with restrictions
Although this study was not conducted to GLP, the test
parameters used were based on a sound scientific design.

09-AUG-2000 (15)

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6. References

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Date: 30-OCT-2001 ID: 68953-84-4

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# 101-72-4

# p-Phenylenediamine, N-Isopropyl-N'-Phenyl-

# 2. PHYSICAL-CHEMICAL DATA

\*2.1 MELTING POINT

Value: 75-80 °C

Decomposition: Yes [] No [X] Ambiguous [] Sublimation: Yes [] No [X] Ambiguous []

Method: FF83.9-1 Initial and Final Melting Point of Organic Compounds

1996

GLP: Yes [X] No [] ? [] Remarks: Capillary Method

Reference: ASTM D-1519 / Flexsys Physical Methods of Analysis

\*2.2 BOILING POINT

Value: 161 °C Pressure: at 1 mm Hg

Decomposition: Yes [] No [X] Ambiguous []

Method: Not listed

GLP: Yes [ ] No [ ] ? [X]

Remarks:

Reference: Monsanto Toxicology Profile of Santoflex IP, 1990

†2.3 DENSITY (relative density)

Type: Bulk density []; Density [X]; Relative Density []

Value: 1.180 Temperature: 20 °C

Method: FF97.8-1 Flexsys Standard Method 1997

GLP: Yes [X] No [] ? []

Remarks: Density of solids by displacement Reference: Flexsys Physical Methods of Analysis

\*2.4 VAPOUR PRESSURE

Value: 0.00343 mm Hg

Temperature: 90 °C

Method: calculated [ ]; measured [ X ]

Not listed

GLP: Yes [ ] No [ ] ? [X]

Remarks:

Reference: Monsanto Toxicology Profile of Santoflex IP, 1990

\*2.5 PARTITION COEFFICIENT log<sub>10</sub>P<sub>ow</sub>

Log Pow: 3.28 Log P
Temperature: Not Determined

Method: calculated [X]; measured []

SRC LogKow (KowWin) Program 1995

GLP: Yes [] No [X] ? []

Remarks:

Reference: Meylan, W.M. and. P.H. Howard, 1995 J. Pharm. Sci. 84: 83-92

# \*2.6 WATER SOLUBILITY

Results:

<b>A.</b>	Solubility				
	Value:	15 ppm			
	Temperature:	25 °C			
	Description:	Miscible []; Of very high solubility [];			
		Of high solubility []; Soluble []; Slightly soluble [];			
	Method:	Of low solubility []; Of very low solubility [X]; Not soluble []			
	GLP:	Saturated Solution / Solvent Extraction / GC.Analysis Yes [ ] No [ ] ? [X]			
	Remarks:	CH2Cl2 solvent, 100% recovery at 1 ppm. Equilibrated w/out			
	Remarks.	light.			
	Reference:	Monsanto ES-78-SS-20, Environmental Sciences, 1978			
В.	pH Value, pKa Value				
	pH Value:	Not Applicable			
	pKa value	5.1 at 25°C			
	Method:	Estimated			
	GLP:	Yes [ ] No [ ] ? [X]			
	Remarks:	Value indicates that this compound will exist only slightly in the cation form			
	Reference:	HSDB database 101-72-4, SRC, University of Georgia SPARC			
	Reference.	SPARC On-Line Calculator			
2.11	OXIDISING PROPERTIES				
	Results:	Maximum burning rate equal or higher than reference mixture[];			
		Vigorous reaction in preliminary test [ ];			
		No oxidising properties [ ]; Other [ ]			
	Method:				
	GLP:	Yes [ ] No [ ] ? [ ]			
	Remarks:				
	Reference:				
†2.12	OXIDATION: REDUCTION POTENTIAL				
	Value:	mV			
	Method:	** ** ** ** **			
	GLP:	Yes [] No [] ? []			
	Remarks:				
	Reference:				
2.13	ADDITIONAL DATA				
A.	Partition co-efficient between soil/sediment and water (Kd)				
	Value:				
	Method:	X			
	GLP:	Yes [ ] No [ ] ? [ ]			
	Remarks: Reference:				
В.	Other data				
	·				

Henry's Law Constant = 1.4 x 10(-9) atm-cu m/mole

Remarks: Fragment Constant Estimation method. Volitazation from moist

soil surfaces is not expected to be an important fate process.

Reference: HSDB – Lyman, W.J. et. al. Handbook of Chemical Property

Estimation Methods, 1990

# 3. ENVIRONMENTAL FATE AND PATHWAYS

*3.1.1	PHOTODEGRADATION				
	Type:	Air [X]; Water [ ]; Soil [ ]; Other [ ]			
	Light source:	Sunlight [ ]; Xenon lamp [ ]; Other [ ]			
	Light spectrum:				
		(based on intensity of sunlight)			
	Spectrum of substance				
	Concentration of Substance:				
	Temperature:	°C			
	Direct photolysis:				
	Half life:				
	Degradation:	(exposure time)			
	Quantum yield:				
	Indirect Photolysis:				
	Type of sensitizer:	ОН			
	Concentration of sensitizer: 1560000 molecule/. cm <sup>3</sup>				
	Rate constant (radical): 218.3766 E-12 cm <sup>3</sup> /molecule*sec				
	Degradation:	50% at 0.588 Hrs			
	Method:	calculated [X]; AOP Program (v1.89)			
		measured [ ]			
	GLP:	Yes [ ] No [ <b>X</b> ] ? [ ]			
	Test substance:	. molecular structure , purity:			
	Remarks:				
	Reliability:	(2) valid with restrictions			
		Accepted calculation method			
	Reference:	Meylan W. and Howard P. (1999) EPIWin Modeling Program.			
		Syracuse Research Corporation. Environmental Science Center,			

# \*3.1.2 STABILITY IN WATER

Type: Abiotic (hydrolysis) [X]; biotic (sediment)[]

Half life: Not Determined

Degradation: 99% at pH 7.0 at 25 °C after 24 Hours

Method: Phase I Hydrolysis Study / ID of Hydrolysis Products

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex IP purple solid Lot # ND02-740, purity: >95%

Remarks: Rapid hydrolysis to benzoquinoneimine-N-phenyl and 4-hydroxy-

diphenylamine. No starting material was detected by GC analysis

6225 Running Ridge Road, North Syracuse, NY 13212-2510.

after 7 days.

Reference: Monsanto ABC-32301, Analytical Bio-Chemistry Labs, 1986

# \*3.2 MONITORING DATA (ENVIRONMENTAL)

Type of Measurement: Background [ ]; At contaminated site [ ]; Other [ ]

Media:

Results: Remarks: Reference:

# 3.3 TRANSPORT AND DISTRIBUTION BETWEEN ENVIRONMENTAL COMPARTMENTS INCLUDING ESTIMATED ENVIRONMENTAL CONCENTRATIONS AND DISTRIBUTION PATHWAYS

# \*3.3.1 TRANSPORT

Type: Adsorption []; Volatility []; Other []

Media: Method: Results: Remarks: Reference:

#### \*3.3.2 THEORETICAL DISTRIBUTION (FUGACITY CALCULATION)

Media: Air-biota []; Air-biota-sediment-soil-water []; Soil-biota [];

Water-air []; Water-biota []; Water-soil []; Other []

Method: Fugacity level I [ ]; Fugacity level II [ ]; Fugacity level III [X];

Fugacity level IV [ ]; Other (calculation) [ ]; Other

(measurement)[ ]

Results: Concentration Half-Life Emissions Fugacity (percent) (hr) (kg/hr) (atm)

1.18 1000 4.69e-013 Air 0.0158 900 1000 1.97e-014 Water 22.4 Soil 76.9 900 1000 3.94e-014 Sediment 0.68 3.6e + 0031.51e-014

Rea	ction 2	Advection	Reaction	Advection
(kg	/hr)	(kg/hr)	(percent)	(percent)
Air	257	4.36	8.57	0.145
Water	1478	620	15.9	20.7
Soil	1.64e+003	3 0	54.6	0
Sediment	3.62	0.376	0.121	0.0125

Persistence Time: 922 hr Reaction Time: 1.16e+003 hr Advection Time: 4.42e+003 hr

Percent Reacted: 79.2 Percent Advected: 20.8

Remarks:

Reliability: (2) valid with restrictions

Accepted calculation method

Reference: Meylan W. and Howard P. (1999) EPIWin Modeling Program.

Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.

# \*3.5 BIODEGRADATION

Type: aerobic [ X ]; anaerobic [ ] Inoculum: adapted [ X ]; non-adapted [ ];

Concentration of the chemical: 1002 ug/l. related to COD []; DOC []; test substance [X] Medium: water [X]; water-sediment []; soil []; sewage treatment []

Degradation: 50% after 2.5 Hours

90 % after 3.5 Hours

98% after 22 Hours

Results: readily biodeg. [X]; inherently biodeg. [V]; under test condition

no biodegradation observed [ ], other [ ]

Method: Natural Water Die-Away Test, Dixon, Hicks and Michael, 1981

GLP: Yes [ X ] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex IP purple solid Lot# N76-7433, purity:>95%.

Remarks: Tests run in Mississippi River Water and purified water. The

short half-lives in both systems suggest that the compound should

not persist in natural aquatic environments.

Reference: Monsanto ES-81-SS-53, MIC Environmental Sciences, 1981

# 4. <u>ECOTOXICITY</u>

# \*4.1 ACUTE/PROLONGED TOXICITY TO FISH

Type of test: static [X]; semi-static []; flow-through []; other) []

open-system [ ]; closed-system [  $\mathbf{X}$  ]

Species: <u>Salmo gairdneri</u> (Rainbow Trout)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 0.62 mg/l

 $LC_{50}$  (48h) = 0.38 mg/l  $LC_{50}$  (72h) = Not reported  $LC_{50}$  (96h) = 0.34 mg/l NOEC = 0.18 mg/l LOEC = 0.24 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: Santoflex IP dark solid, Lot#NO12-002, purity: >95%

Remarks: Stock solutions prepared in reagent-grade acetone. Water quality

parameters of temperature, dissolved oxygen and pH monitored throughout test. Observations and mortality counts were made

every 24 hours.

Reference: Monsanto BN-76-255, EG&G Bionomics, 1977

Type of test: static [X]; semi-static []; flow-through []; other) []

open-system [ ]; closed-system [X]

Species: Lepomis machrochirus (Bluegill Sunfish)

Exposure period: 96 Hours

Results:  $LC_{50}$  (24h) = 0.48 mg/l

 $LC_{50}$  (48h) = 0.43 mg/l  $LC_{50}$  (72h) = Not reported  $LC_{50}$  (96h) = 0.43 mg/l NOEC = 0.24 mg/l LOEC = 0.32 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: Santoflex IP dark solid, Lot# NO12-002, purity: >95%

Remarks: Stock solutions prepared in reagent-grade acetone. Water quality

parameters of temperature, dissolved oxygen and pH monitored

throughout test. Observations and mortality counts were made

every 24 hours

Reference: Monsanto BN-76-255, EG&G Bionomics, 1977

Type of test: static []; semi-static []; flow-through [X]; other []

open-system [ ]; closed-system [ X ]

Species: <u>Pimephales promelas</u> (Fathead Minnows)

Exposure period: 14 days

Results:  $LC_{50}$  (24h) = 1.80 mg/l

 $\begin{array}{l} LC_{50} \ (192h) = 0.28 \ mg/l \\ LC_{50} \ (240h) = 0.21 \ mg/l \\ LC_{50} \ (336h) = 0.09 \ mg/l \end{array}$ 

Analytical monitoring: Yes [X] No [] ? []

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [X] No [] ? [] Klimisch 1

Test substance: Santoflex IP dark solid rec'd 4/25/78, purity: >95%

Remarks: Stock solutions prepared in reagent-grade acetone. Water quality

parameters of temperature, dissolved oxygen, ammonia and pH monitored throughout test. Although the goal of the study was to determine a lethal threshold concentration of the test substance, the results indicated that this was not reached at 14 days. In addition, the test substance appeared to exhibit cumulative

toxicity to the fish under test conditions.

Reference: Monsanto AB78-120B, Analytical Bio-Chemistry Labs, 1979

Type of test: static [X]; semi-static []; flow-through []; other) []

open-system [ ]; closed-system [X]

Species: Paratanytarsus parthenogenetica (Midge)

Exposure period: 48 Hours

Results:  $LC_{50}$  (24h) = 29 mg/l

 $LC_{50}$  (48h) = 23 mg/l NOEC = Not Observed

LOEC = 10 mg/l (lowest concentration tested)

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975) and Gettings and Adams, Method for Conducting Acute Toxicity Tests with Midge

1980

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex IP #1803025-C), purity: >95%

Remarks: Stock solutions prepared in reagent-grade acetone. Water quality

parameters of temperature, dissolved oxygen, ammonia and pH

monitored throughout test.

Reference: Monsanto 9AB981013, Analytical Bio-Chemistry Labs, 1981

# 4.2 ACUTE TOXICITY TO AQUATIC INVERTEBRATES

#### \*A. Daphnia

Type of test: static [ X ]; semi-static [ ]; flow-through [ ]; other [ ];

open-system [ ]; closed-system [ X ]

Species: <u>Daphnia magna</u>
Exposure period: 48 Hours

Results:  $EC_{50}$  (24h) = 2.8 mg/l

 $EC_{50}$  (48h) = 1.1 mg/l NOEC = 0.56 mg/l

Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians (1975)

GLP: Yes [ ] No [ ] ? [X] Klimisch 2

Test substance: Santoflex IP purple flakes Lot #676-7433, purity: >95%

Remarks: Acetone used to prepare stock solutions. Initial range-finding

experiment run to determine appropriate concentrations for final experiment. Water quality parameters of dissolved oxygen, pH, hardness, temperature and alkalinity monitored throughout the

test.

Reference: Monsanto AB-78-120, Analytical Bio-Chemistry Labs, 1978

# \*4.3 TOXICITY TO AQUATIC PLANTS, e.g. algae

Species: <u>Selanastrum capricurnutum</u> (Freshwater alga)
Endpoint: Biomass [ ]; Growth rate [ **X** ]; Other [ ]

Exposure period: 96 Hours

Results:  $EC_{50}$  (96h) = 0.4 ppm for a chlorophyll, 0.5 ppm for cell numbers

NOEC = <0.1 ppm LOEC = Not Determined Analytical monitoring: Yes [X] No [ ] ? [ ]

Method: US EPA Algal Test Procedure: Bottle Test, 1971

open-system []; closed-system [X]

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex IP #BN-78-1384325, purity: >95%

Remarks: Both a chlorophyll and cell numbers measured to confirm results.

Stock solutions prepared in acetone; acetone also used as solvent control Concentrations of test article determined by preliminary

range-finding experiment.

Reference: Monsanto BN-78-1384325, EG&G Bionomics, 1978

# 5. <u>TOXICITY</u>

# \*5.1 ACUTE TOXICITY

# 5.1.1 ACUTE ORAL TOXICITY

Type:  $LD_0[]; LD_{100}[]; LD_{50}[X]; LDL_0[]; Other[]$ 

Species/strain: Sprague-Dawley Albino Rats

Value: 900 mg/kg b.w.:

Discriminating dose: 1000 mg/kg/bw

Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [ X ] Klimisch 2
Test substance: Santoflex IP Lot# NO12-002, purity: <95%

Remarks: The test article was administered to groups of male and female

rats by oral gavage as a 20% suspension in corn oil vehicle. Dose levels were 631, 794, 1000 or 1260 mg/kg/bw. Clinical signs of toxicity were reduced appetite and activity – three to five days in

survivors – followed by increasing weakness, collapse and death. Most deaths occurred within two days. Gross autopsy findings on decedents included lung hyperemia, slight liver discoloration and acute gastrointestinal inflammation. Survivors were sacrificed after a two-week recovery period. All viscera examined appeared normal in these animals.

Reference: Monsanto Y-73-287, Younger Laboratories, 1974

# 5.1.2 ACUTE INHALATION TOXICITY

Type:  $LC_0[]; LC_{100}[]; LC_{50}[]; LCL_0[]; Other[]$ 

Species/strain: Exposure time: Value:

Method:

GLP: Yes [ ] No [ ] ? [ ]

Test substance: ...., purity:....

Remarks: Reference:

#### 5.1.3 ACUTE DERMAL TOXICITY

Type:  $LD_0[]$ ;  $LD_{100}[]$ ;  $LD_{50}[X]$ ;  $LDL_0[]$ ; Other[]

Species/strain: New Zealand Albino Rabbits

Value: >7940 mg/kg b.w.
Method: Defined Lethal Dose

GLP: Yes [ ] No [ ] ? [X] Klimisch 2
Test substance: Santoflex IP Lot #NO12-002, purity: >95%

Remarks: The test article was applied to the shaved skin of groups of male

and female rabbits for 24-hours as a 40% suspension in corn oil. Doses wer either 5010 or 7940 mg/kg/bw. All animals survived until sacrifice. Clinical signs of toxicity were limited to reduced appetite and activity for three to five days. Following a two-week recovery period, the animals were sacrificed. All viscera

examined appeared normal in all animals.

Reference: Monsanto Y-73-287, Younger Laboratories, 1974

#### \*5.4 REPEATED DOSE TOXICITY

Species/strain: Sprague-Dawley Albino Rats

Sex: Female [ ]; Male [ ]; Male/Female [ X ]; No data [ ]

Route of Administration: Oral/Dietary

Exposure period: 30 Days
Frequency of treatment: Daily
Post exposure observation period:

Dose: 0, 500, 1000, 1750 or 2500 ppm Control group: Yes [X]; No []; No data [];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

NOEL: 500 ppm LOEL: 1000 ppm

Results: In a 30-day range-finding study that preceded a 90-day study,

the test substance was administered orally, via dietary admixture, to groups of male and female rats (5/sex/group). Control animals received the standard laboratory diet. Physical observations, body weight and food consumption measurements were

performed on all animals pretest and at selected intervals during the study. Hematology and chemistry determinations were performed on all animals at study termination. There were no mortalities during the course of the study. After four weeks of treatment, all animals were sacrificed, selected organs were weighed, and organ/body weight ratios were calculated. Complete postmortem examinations were conducted on all animals. Differences from control in body weight gain, hematological effects, elevations in total serum protein and increased liver and spleen weights were noted in animals dosed at 1000 ppm and above. There were no significant differences in findings between control groups animals and those dosed at 500 ppm that were attributed to the test article.

Dunnett, C.W., A Multiple Comparison Procedure for Method:

Comparing Several Treatments with a Control, Jour. Am. Stat.

Assoc. 50: 1096-1121, 1955

GLP: Yes [X] No [] ? [] Klimisch 1 Santoflex IP Lot# 7J111, purity: 97.2% Test substance: Monsanto BD-88-74, Bio/dynamics Inc. 1988 Reference:

Species/strain: Sprague-Dawley Albino Rats

Sex: Female [ ]; Male [ ]; Male/Female [X]; No data [ ]

Route of Administration: Oral/Dietary

Exposure period: 90 Days Frequency of treatment: Daily Post exposure observation period:

0, 180, 360 or 720 ppm

Control group: Yes [ ]; No [ ]; No data [ ];

Concurrent no treatment [X]; Concurrent vehicle []; Historical []

180 ppm for males, Not determined for females NOEL: LOEL: 360 ppm for males, 180 ppm for females

Results:

The test substance was administered orally, via dietary admixture, to groups of male and female rats (10/sex/group). Control animals received the standard laboratory diet. Physical observations, body weight and food consumption measurements were performed on all animals pretest and at selected intervals during the study. Hematology and chemistry determinations were performed on all animals at Months 1.5 and 3. One high-dose and one mid-dose female were found dead on test day 93 following collection of terminal blood samples. The cause of death was attributed to the stress of bleeding and not to the administration of the test article. There were no other mortalities during the course of the study. After three months of treatment, all animals were sacrificed, selected organs were weighed, and organ/body and organ/brain weight ratios were calculated. Complete postmortem examinations were conducted on all animals. Histopathological evaluation of selected tissues was performed on all control and high-dose animals. The lungs, spleen, liver and kidneys were examined microscopically for all animals in all groups. Mean body weights and mean body weight gains were slightly reduced (2-4%) in males at 750 ppm.

Treatment-related findings were observed in several hematology parameters in the males and/or females at dose levels of 360 and 720 ppm. Parameters affected included reduced hemoglobin concentrations and hematocrit values at Week 6, reduced hemoglobin concentration in 720 ppm females at Week 13, elevated platelet counts in males at Week 6, and reduced mean erythrocyte counts in females at Week 6 and in high-dose females only at Week 13. The NOEL for hematology data was set at 180 ppm for both sexes. Differences in clinical chemistry parameters were noted in all mid- to high-dose animals. Mean liver weights, liver-to-body-weight and liver-to-brain-weight ratios were increased in 360 and 720 males, and in all treated females. There were no treatment-related findings noted in mortality, physical observations, opthalmology, food consumption or gross or

microscopic pathology in any dose/sex group.

Method: OECD Guidelines for Testing of Chemicals, Section 453, 1981

and US EPA TSCA Section 4(a) Test Rules, 1982

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex IP Lot# 7J111, purity: 97.2%

Reference: Monsanto BD-88-389, Bio/dynamics, Inc. 1990

#### \*5.5 GENETIC TOXICITY IN VITRO

## A. BACTERIAL TEST

Type: Bacterial Reverse Mutation - Ames

System of testing: TA-98, TA-100, TA-1535, TA-1537, TA-1538 Concentration: 0.1, 1,10, 100 and 500 micrograms/plate

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 500 ug/plate

Without metabolic activation: 500 ug/plate

Precipitation conc: Not determined

Genotoxic effects: + ? -

With metabolic activation: [ ] [ ] [X] Without metabolic activation: [ ] [ ] [X]

Method: Ames Plate Test (Overlay method) 1975; OECD 471 equivalent

GLP: Yes [] No [] ? [X] Klimisch 2
Test substance: Santoflex IP 11-02-76, purity: >95%

Remarks: Stock solutions prepared in DMSO. No evidence of mutagenic

activity in any assay conducted with or without activation using

the S-9 homogenate from Arochlor-induced rat livers.

Reference: Monsanto BIO-76-226, Litton Bionetics, 1976

Type: Bacterial Reverse Mutation - Ames
System of testing: TA-98, TA-100, TA-1535, TA-1537

Concentration: 0.2, 0.8, 4, 20, 60 and 200 micrograms/plate

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 200 ug/plate

Without metabolic activation: 200 ug/plate

Precipitation conc: Insoluble at 1 mg/plate and above

Genotoxic effects: + ? -

With metabolic activation: [] [] [X] Without metabolic activation: [] [] [X]

Method: Ames Plate Test (Overlay method) 1975; OECD 471 equivalent

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex IP Lot# ND02-740, purity: 92-99%

Remarks: Stock solutions prepared in DMSO. No evidence of mutagenic

activity in any assay conducted with or without activation using

the S-9 homogenate from Arochlor-induced rat livers.

Reference: Monsanto ML-85-243, Environmental Health Labs, 1986

# B. NON-BACTERIAL IN VITRO TEST

Type: Mammalian Cell Gene Forward Mutation Assay

System of testing: L5178Y Mouse Lymphoma cells

Concentration: 0.156, 0.313, 0.625, 1.250, 2.500 (without activation)

0.625, 1.250, 2.500, 5.000 and 10.000 (with activation)

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: With metabolic activation: 10.0 ug/ml

Without metabolic activation: 2.5 ug/ml

Precipitation conc: >1 mg/ml

Genotoxic effects: + ? -

With metabolic activation: [][][X] Without metabolic activation: [][][X]

Method: Clive and Spector, Mutation Research 31:17-29 (1975)

GLP: Yes [] No [] ? [X] Klimisch 2
Test substance: Santoflex IP flakes Lot # N76-7433, purity 97%

Remarks: The test article was evaluated for specific locus forward mutation

in the L5178Y Thymidine Kinase (TK) mouse lymphoma cell assay. Stock solutions were prepared in DMSO. DMSO was used as the negative control. EMS was used as the positive control without activation and DMN was used as the positive control with activation. The test article was found to be negative

Reference: Monsanto BIO-78-224 Litton Bionetics, 1978

Type: <u>In vitro</u> Unscheduled DNA Synthesis (UDS)

System of testing: Primary rat hepatocyte cultures (Fischer-344 strain) Concentration: 0.01, 0.05, 0.1, 0.5, 1, 3, 5, 10, 50, 100, 1000 ug/ml

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Cytotoxicity conc: Preliminary Assay: 5 ug/ml

Replicate Assay: 3 ug/ml

Precipitation conc: Separation/sticking to sides of tube noted at 100 ug/ml and above

Genotoxic effects: + ? -

[][][X]

Method: Williams, G.M., Detection of Chemical Carcinogens by

Unscheduled DNA Synthesis in Rat Liver Primary Cell Cultures,

Cancer Research <u>37</u>, pp. 1845-1851 (1977)

GLP: Yes [X] No [ ] ? [ ] Klimisch 1

Test substance: Santoflex IP flakes Lot# ND02-740, purity 92-97%

Remarks: Acetone (1%) used as solvent and diluent. Primary rat liver cell

cultures derived from the livers of two adult male rats. The positive control was 2-AAF, the solvent control was acetone in the preliminary assay and DMSO in the replicate assay. The percentage of cells in repair was calculated as the percentage of cells with at least 5 net grains/nucleus. 150 cells were scored for each concentration reported for each experiment. The net grain counts were negative at each concentration of the test compound, in the solvent control and in the medium control, in contrast to the strong positive response produced by the positive control 2-AAF in both experiments. These results indicate that Santoflex IP is not a genotoxic agent under the conditions of the in vitro rat

hepatocyte DNA repair assay.

Reference: Monsanto SR-85-251, SRI International, 1986

Type: CHO/HGPRT Forward Gene Mutation Assay System of testing: Cultured Chinese hamster ovary (CHO) cells

Concentration: 2, 5, 10, 15 and 30 ug/ml

Metabolic activation: With []; Without []; With and Without [X]; No data []

Results:

Method:

Cytotoxicity conc: With metabolic activation: 30 ug/ml

Without metabolic activation: 10 ug/ml

Precipitation conc: Not Determined

Genotoxic effects: + ? -

With metabolic activation: [] [] [X]
Without metabolic activation: [] [] [X]
CHO/HGPRT Mutation Assay (1979) Hsie, et.al.

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex IP Lot# N002-740, purity: 92-99%
Remarks: The mutagenic potential of Santoflex IP was t

The mutagenic potential of Santoflex IP was tested in CHO cells for ability to induce forward mutation at the HGPRT gene locus. A range-finding cytotoxicity study preceded a dose-response mutagenicity experiment using different levels of Arochlor1254 rat liver homogenate (S9) concentrations, followed by a confirmatory dose-response mutagenicity experiment. The compound was tested at S9 concentrations up to a cytotoxic dose of 30 ug/ml. No statistically significant mutagenicity was observed in the two separate experiments. Therefore, the test substance was not considered to be mutagenic in CHO cells under

the experimental conditions.

Reference: Monsanto ML-85-221, Environmental Health Labs, 1986

# \* 5.6 GENETIC TOXICITY IN VIVO

Type:

Species/strain:

Sex: Female [ ]; Male [ ]; Male/Female [ ]; No data [ ]

Route of Administration:

Exposure period:

Doses: Results:

Effect on mitotic index or P/N ratio: + ? -Genotoxic effects: [11111]Method: GLP: Yes [ ] No [ ] ? [ ] Test substance: ...., purity: ..... Remarks: Reference: TOXICITY TO REPRODUCTION Fertility []; One-generation study []; Two-generation study []; Other [ ] Species/strain: Sex: Female [ ]; Male [ ]; Male/Female [ ]; No data [ ] Route of Administration: Exposure period: Frequency of treatment: Post exposure observation period: Premating exposure period: male: . . . . , female: Duration of the test: Doses: Control group: Yes [ ]; No [ ]; No data [ ]; Concurrent no treatment []; Concurrent vehicle []; Historical [] **NOEL Parental:** NOEL F1 Offspring: NOEL F2 Offspring:

Results:

General parental toxicity Toxicity to offspring:

Method:

\*5.8

GLP: Yes [] No [] ? [] Test substance: ....., purity: .....

Remarks: Reference:

#### \*5.9 DEVELOPMENTAL TOXICITY/ TERATOGENICITY

Species/strain: Sprague-Dawley CD Rats

Sex: Female [X]; Male []; Male/Female []; No data []

Route of Administration: Oral gavage

Duration of the test: 20 days from mating to C-section

Exposure period: Day 6-15 of gestation

Frequency of treatment: Daily, as a single oral dose at a volume of 5 ml/kg

Doses: 0, 12.5, 62.5 and 125 mg/kg/bw Yes [ X ]; No [ ]; No data [ ]; Control group:

Concurrent no treatment []; Concurrent vehicle [X]; Historical []

NOEL Maternal Toxicity: 62.5 mg/kg NOEL teratogenicity: 62.5 mg/kg

Results: The test substance was administered to groups of 24 pregnant rats

> during the period of embryo organogenesis. The vehicle was Polyethylene Glycol 400, and dose levels were 0, 12.5, 62.5 or

125 mg/kg/bw.

Maternal general toxicity: High-dose rats exhibited slight maternal toxicity as evidenced by a reduction in food intake, predosing salivation and soft, dark feces. There were no effects on body weight. All animals survived to sacrifice. There were no treatment-related macroscopic findings at necropsy for any dose level

Pregnancy/litter data: There were no treatment-related effects on uterine/implantation.

Foetal data: At 125 mg/kg there were statistically significant effects on the incidence of skeletal findings. Effects included an increased incidence of irregularly and incompletely ossified cranial and facial bones, and increased incidence of no ossification of hyoid, unilateral/bilateral wavy ribs, and semi-bipartite vertebral centra. At 62.5 mg/kg, there was a statistically significant increase in incomplete ossification of more than one cranial bone. At 12.5 mg/kg, there was a statistically significant increase in the incomplete ossification of more than one facial bone that was not considered to be treatment-related.

Method: OECD 59B (1982)

GLP: Yes [X] No []? [] Klimisch 1
Test substance: Santoflex IP dark flakes, Lot#2F054, purity: 97%

Remarks: No deviations from protocol noted.

Reference: Monsanto SP-93-46, SafePharm Laboratories 1994

#### 5.10 OTHER RELEVANT INFORMATION

## A. Specific toxicities

Type: Immunotoxicity

Repeat Insult Patch Test

Results: Santoflex IP, 50% w/v in Dimethylphthalate, was applied to the

upper arm of 50 human volunteers using a linteen disk moistened with the test material. The patch was kept in place for 24 hours before removal and grading of gross skin changes on a scale of 0-4. After a 24-hour rest period, the test material was reapplied. This cycle was repeated every Monday, Wednesday and Friday, with a 48-hour rest period over weekends. After the 15<sup>th</sup> application, the volunteers rested two weeks before the challenge application.

Application #1: Score 0/50 Applications #2-15: Score 10/50 Challenge: Score 11/50

Remarks: Under the test conditions, 11/50 or 22% of the volunteers showed

sensitization responses. Those 11 persons were also subjected to a supplementary challenge using Santoflex 13 (6PPD). No subject showed any indication of cross-sensitization from one

PPD rubber chemical material to another.

Reference: Monsanto SH-76-7, Product Investigations, Inc. 1976

Type: Immunotoxicity

Modified Draize Skin Sensitization Study on Human Volunteers

Results: The study was performed over a 6-week period on 82 human

volunteers using Santoflex IP, 1%, in petrolatum. During the first three weeks, patches moistened with the test material were applied to the arms at the same site at the rate of three times/week. Following a rest period, a challenge application was made to a different site. Results for irritation and sensitization were scored on a scale of 0-4. 12 of 82 test subjects were deemed to be sensitized, for a rate of 14.6%

Reference: Monsanto MA-78-92, 1978

# B. Toxicodynamics, toxicokinetics

Type:

Results:

Remarks:

References:

## \* 5.11 EXPERIENCE WITH HUMAN EXPOSURE

Results:

Remarks:

Reference:

#### 6. REFERENCES

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- 14. Monsanto AB-78-120, Acute Toxicity of Santoflex IP to <u>Daphnia magna</u>, Analytical Bio-Chemistry Laboratories, Inc. August 25, 1978
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IUCLID

Data Set

Existing Chemical ID: 793-24-8 CAS No. 793-24-8

EINECS Name N-1,3-dimethylbutyl-N'-phenyl-p-phenylenediamine

EINECS No. 212-344-0

TSCA Name 1,4-Benzenediamine, N-(1,3-dimethylbutyl)-N'-phenyl-

Molecular Formula C18H24N2

Producer Related Part

Company:

Creation date: 23-SEP-1999

Substance Related Part

Company:

Creation date: 23-SEP-1999

Memo: RAPA PPD category

Printing date: 20-NOV-2001

Revision date:

Date of last Update: 20-NOV-2001

Number of Pages: 57

Chapter (profile): Chapter: 1, 2, 3, 4, 5, 7

Reliability (profile): Reliability: without reliability, 1, 2, 3, 4

Flags (profile): Flags: without flag, confidential, non confidential, WGK

(DE), TA-Luft (DE), Material Safety Dataset, Risk

Assessment, Directive 67/548/EEC, SIDS

Date: 20-NOV-2001

1. General Information ID: 793-24-8

#### 1.0.1 OECD and Company Information

Type: lead organisation

Name: American Chemistry Council (formerly Chemical Manufacturers

Association), Rubber and Plastic Additives Panel

Street: 1300 Wilson Boulevard Town: 22209 Arlington, VA

Country: United States Phone: 703-741-5600 Telefax: 703-741-6091

20-NOV-2001

Type: cooperating company
Name: Bayer Corporation
Country: United States

20-NOV-2001

Type: cooperating company

Name: Ciba Specialty Chemicals Corporation

Country: United States

20-NOV-2001

Type: cooperating company Name: Crompton Corporation

Country: United States

20-NOV-2001

Type: cooperating company Name: Flexsys America L.P.

Country: United States

20-NOV-2001

Type: cooperating company

Name: Noveon, Inc. (formerly BF Goodrich)

Country: United States

20-NOV-2001

Type: cooperating company

Name: R.T. Vanderbilt Company, Inc.

Country: United States

20-NOV-2001

Type: cooperating company

Name: The Goodyear Tire & Rubber Company

Country: United States

20-NOV-2001

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Date: 20-NOV-2001

1. General Information

Date: 793-24-8

Type: cooperating company
Name: The Lubrizol Corporation

Country: United States

20-NOV-2001

Type: cooperating company Name: UOP, LLC.

Name: UOP, LLC. Country: United States

20-NOV-2001

1.0.2 Location of Production Site

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1.0.3 Identity of Recipients

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1.1 General Substance Information

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1.1.0 Details on Template

\_

1.1.1 Spectra

\_

1.2 Synonyms

-

1.3 Impurities

\_

1.4 Additives

\_

1.5 Quantity

\_

1.6.1 Labelling

\_

1.6.2 Classification

\_

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Date: 20-NOV-2001

1. General Information

ID: 793-24-8

1.7 Use Pattern

-

1.7.1 Technology Production/Use

\_

1.8 Occupational Exposure Limit Values

\_

1.9 Source of Exposure

-

1.10.1 Recommendations/Precautionary Measures

-

1.10.2 Emergency Measures

\_

1.11 Packaging

\_

1.12 Possib. of Rendering Subst. Harmless

\_

1.13 Statements Concerning Waste

\_

1.14.1 Water Pollution

-

1.14.2 Major Accident Hazards

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1.14.3 Air Pollution

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1.15 Additional Remarks

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1.16 Last Literature Search

-

- 3/57 -

Date: 20-NOV-2001

1. General Information

ID: 793-24-8

1.17 Reviews

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1.18 Listings e.g. Chemical Inventories

-

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2. Physico-chemical Data 1D. 793-24-0

2.1 Melting Point

Value: 45 degree C

Decomposition: no Sublimation: no

Method: other: FF83.9-1 Initial and Final Melting Point of Organic

Compounds.

Year: 1996 GLP: yes

Testsubstance: other TS: CAS# 793-24-8

Remark: Capillary method

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (1)

Value: 50 degree C

Method: other: Handbook value

GLP: no data

Testsubstance: other TS: CAS# 793-24-8
Reliability: (2) valid with restrictions

Data from Handbook or collection of data

Flag: Critical study for SIDS endpoint

20-NOV-2001 (2)

Value: 45 - 48 degree C Source: Bayer AG Leverkusen

20-NOV-2001 (3)

2.2 Boiling Point

Value: 230 degree C at 13.3 hPa Source: Bayer AG Leverkusen

28-SEP-1992 (3)

2.3 Density

Type: relative density Value: 1 at 15 degree C

Method: other: FF97.8-1 Flexsys Standard Method

Year: 1997 GLP: yes

Testsubstance: other TS: CAS# 793-24-8

Remark: Density of solids by displacement Flag: Critical study for SIDS endpoint

20-NOV-2001 (4)

Type:

Value: 1.02 g/cm3 at 20 degree C Source: Bayer AG Leverkusen

28-SEP-1992 (3)

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## 2. Physico-chemical Data

Type:

Value: .995 g/cm3 at 50 degree C Source: Bayer AG Leverkusen

20-JUN-1997 (5)

relative density Type: 1 at 60 degree C Value:

MonsantoBayer AG Leverkusen Source:

26-MAY-1994

#### 2.3.1 Granulometry

#### 2.4 Vapour Pressure

Value: 8.7 hPa at 200 degree C Source: Bayer AG Leverkusen

28-SEP-1992 (3)

Value: 93 hPa at 300 degree C Bayer AG Leverkusen Source:

28-SEP-1992 (3)

# 2.5 Partition Coefficient

log Pow: 4.68 at 25 degree C

Method: other (calculated): SRC LogKow (KowWin) Program

Year: 1995 GLP: no

Testsubstance: other TS: molecular structure (2) valid with restrictions Reliability: Accepted calculation method

Flaq: Critical study for SIDS endpoint

20-NOV-2001 (6)

log Pow: 5.4

Method: other (calculated): Leo, A.: CLOGP-3.54 MedChem Software 1989.

Daylight, Chemical Information Systems, Claremont,

CA 91711, USA

Year:

Bayer AG Leverkusen Source:

(2) valid with restrictions Reliability:

(7)20-NOV-2001

log Pow: Method: Year:

Remark: pow = 59000 + / - 34000Source: Bayer AG Leverkusen

14-JAN-1993 (8)

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## 2. Physico-chemical Data

#### 2.6.1 Water Solubility

Value: 1.1 other: ppm at 23 degree C

Qualitative: not soluble

Method: other: Saturated Solution / Solvent Extraction / GC.Analysis

no data GLP:

Testsubstance: other TS: CAS# 793-24-8

CH2Cl2 solvent, 96% recovery at 1 ppm. Equilibrated w/out Remark:

light.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Critical study for SIDS endpoint Flaq:

20-NOV-2001 (9)(10)

Value: ca. 1 mg/l at 50 degree C

other: modified OECD Guideline 105 "Water solubility-Flask Method:

Method"

Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

20-NOV-2001 (5)

#### 2.6.2 Surface Tension

#### 2.7 Flash Point

200 degree C Value: closed cup Type:

other: DIN 51758 Method:

Year:

Source: Bayer AG Leverkusen

28-SEP-1992 (3)

#### 2.8 Auto Flammability

## 2.9 Flammability

Result:

no information Remark:

Bayer AG Leverkusen Source:

04-FEB-1992

#### 2.10 Explosive Properties

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Date: 20-NOV-2001
2. Physico-chemical Data

ID: 793-24-8

2.11 Oxidizing Properties

-

2.12 Additional Remarks

\_

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Date: 20-NOV-2001 ID: 793-24-8

#### 3. Environmental Fate and Pathways

#### 3.1.1 Photodegradation

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Conc. of sens.: 1560000 molecule/cm3

Rate constant: .0000000002264928 cm3/(molecule \* sec)

Degradation: 50 % after .6 hour(s)

Method: other (calculated): AOP Program (v1.89) Year: 1999 GLP: no

Test substance: other TS: molecular structure Reliability: (2) valid with restrictions Accepted calculation method

Flaq: Critical study for SIDS endpoint

20-NOV-2001 (11)

Type: air INDIRECT PHOTOLYSIS Sensitizer: OH

Method: other (calculated): calculation according to Atkinson

Year: GLP:

Test substance: other TS: CAS# 793-24-8

t1/2 = 1.1 hRemark:

Bayer AG Leverkusen Source:

(2) valid with restrictions Reliability: Accepted calculation method

Flag: Critical study for SIDS endpoint

20-NOV-2001

#### 3.1.2 Stability in Water

abiotic Type:

93 % after Degradation: 24 hour(s)

at pH 70 and 25 degree C

yes Deg. Product:

Method: other: Phase I Hydrolysis Study / ID of Hydrolysis Products

Year: GLP: yes

Test substance: other TS: Purple solid # KD08-281 purity: >95%

Rapid hydrolysis to 4-Hydroxylamine and Remark:

Benzoquinoneimine-N-phenyl.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Critical study for SIDS endpoint Flaq:

20-NOV-2001 (12)

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Date: 20-NOV-2001 ID: 793-24-8

#### 3. Environmental Fate and Pathways

Type: abiotic

Degradation: = 60 % after 25 hour(s)

Method: other: Monsanto Laboratory protocol; see test conditions

GLP: no data Year: 1978

Test substance:

Remark: Degradation data versus time: 0 hour 1 mg/l, 1 hour 0.855

mg/l, 2 hour 0.846 mg/l, 3.5 hour 0.636 mg/l and 25 hour

0.402 mg/l

MonsantoBayer AG Leverkusen Source:

Degradation of test substance in deionized water Test condition:

Reliability: (2) valid with restrictions

20-OCT-1999 (13)

Type: abiotic

t1/2 pH7: = 3 - 4 hour(s) at 24 degree C

Method: other: Monsanto Laboratory protocol; see test conditions

Year: 1993 GLP: yes

Test substance:

Remark: Santoflex 13 is an antiozonant and as such necessarily

> reacts very quickly with oxygen. Therefore, fast oxidation in dilute solutions, where oxygen is readily available, is to be expected. The initial oxidation product is believed

to be quinondiimine, which itself is a very reactive

species. The quinondiimine can hydrolyze or form a polymer by further oxidation giving very complicated mixtures of

products usually involving loss of the alkyl group.

MonsantoBayer AG Leverkusen Source:

Degradation in pH 7 buffered deionized water Test condition:

30-MAY-1994 (14)

## 3.1.3 Stability in Soil

Type: Radiolabel:

Concentration: Cation exch. capac. Microbial biomass: Method:

> Year: GLP:

Test substance:

Remark: no information

Source: Bayer AG Leverkusen

12-JUN-1992

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#### 3. Environmental Fate and Pathways

#### 3.2 Monitoring Data (Environment)

Type of

measurement:

Medium: Method:

Concentration

Remark: no information Bayer AG Leverkusen Source:

06-FEB-1992

#### 3.3.1 Transport between Environmental Compartments

Type: fugacity model level III

Media: other: air, water, soil, sediment

Air (Level I): Water (Level I): Soil (Level I): Biota (L.II/III): Soil (L.II/III):

Method: other: EPIWIN Level III Fugacity Model

Year: 1999

Result: Media Concentration Half-Life Emissions Fugacity (hr) (percent) (kg/hr) (atm) Air 0.0264 1.13 1000 6.66e-013 900 3.36e-014 Water 19.6 1000 900 Soil 68.1 1000 2.84e-015 Sediment 12.2 3.6e+003 0 2.28e-014

Media	Reaction	Advection	Reaction	Advection
	(kg/hr)	(kg/hr)	(percent)	(percent)
Air	457	7.47	15.2	0.249
Water	427	555	14.2	18.5
Soil	1.48e+003	0	49.4	0
Sediment	66.2	6.88	2.21	0.229

Persistence Time: 941 hr Reaction Time: 1.16e+003 hr Advection Time: 4.96e+003 hr

Percent Reacted: 81 Percent Advected: 19

Reliability: (2) valid with restrictions Accepted calculation method

Critical study for SIDS endpoint Flaq:

20-NOV-2001 (11)

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#### 3. Environmental Fate and Pathways

#### 3.3.2 Distribution

air - biota - sediment(s) - soil - water Media: Method: other (calculation): Fugacity Level III

Year: 1999

Result: Concentration Half-Life Emissions

> (kg/hr) (percent) (hr) 0.0264 1.13 1000 Air 19.6 900 1000 Water Soil 900 68.1 1000 Sediment 12.2 3600 0

(2) valid with restrictions Reliability:

21-OCT-1999 (15)

Media: Method: Year:

Remark: Based on the calculated log Pow, transport of the compound

from water to soil/sediment (geoaccumulation) is to be

expected.

Water solubility and vapour pressure indicate that the

transport from water to air is of low relevance.

Bayer AG Leverkusen Source:

21-OCT-1999

#### 3.4 Mode of Degradation in Actual Use

no information Remark:

Source: Bayer AG Leverkusen

06-FEB-1992

- 12/57 -

3. Environmental rate and ratinary

#### 3.5 Biodegradation

Type: aerobic

Inoculum: other: Mississippi River water

Concentration: 1.002 mg/l related to Test substance

Degradation: = 97 % after 22 hour(s)

Result: other: Primary degradation, 96 % primary degradation in

sterile river water and 88 % in deionized water in 22 hours

Testsubstance: 1 hour(s) = 40 % 2 hour(s) = 57 %

3 hour(s) = 67 % 4 hour(s) = 62 % 5 hour(s) = 74 %

Method: other: Natural Water Die-Away in Mississippi River water

Year: GLP: yes

Test substance: other TS: Santoflex 13 Lot# KD-03017, purity: >95%

Remark: Rate of disappearance in

time active sterile deionized Mississippi Mississippi water River water River water 100 % 100 % 100 % 0 hour 100 % 1 hour 60 % 85 % 2 hour 43 % 70 % 88 % 33 % 56 % 86 % 3 hour 49 % 4 hour 38 % 80 % 5 hour 26 % 41 % 65 % 22 hour 3 % 4 % 12 %

Result: 50% degradation after 2.9 hours Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (16)

Type: aerobic

Inoculum: predominantly domestic sewage

Degradation: 13 - 40 % after 28 day

Method: other: Respirometer-Test, ISO DP 9408, EG Directive 79/831/

Annex V, modified MITI Test

Year: GLP: no

Test substance: other TS

Source: Bayer AG Leverkusen Test substance: technical grade 6PPD

20-NOV-2001 (17)

- 13/57 -

Date: 20-NOV-2001 ID: 793-24-8

#### 3. Environmental Fate and Pathways

Type: aerobic

Inoculum: activated sludge

Concentration: 30 mg/l related to Test substance
Degradation: = 7.2 % after 32 day
Result: other: 7.2 % CO2 evolution in 32 days

other: Method similar to Gledhill method listed in U.S.E.P.A. Method:

40 CFR Ch 1 subpart D paragraph 796.3100.

GLP: no data Year:

Test substance:

MonsantoBayer AG Leverkusen Source:

20-NOV-2001 (13)

3.6 BOD5, COD or BOD5/COD Ratio

3.7 Bioaccumulation

#### 3.8 Additional Remarks

Remark: 1.4-Benzenediamine, N-(1.3-dimethylbutyl)-N'-phenyl

decreases the degradation rate of unprotected rubber

(vulcanisate) in water.

Source: Bayer AG Leverkusen

01-DEC-1992 (18)

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#### AQUATIC ORGANISMS

#### 4.1 Acute/Prolonged Toxicity to Fish

Type: static

Species: Salmo gairdneri (Fish, estuary, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: yes

LC50: = .14

Method: other: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians.

Year: 1977 GLP: no data

Test substance: other TS: Santoflex 13 Lot# KD03-017, purity: >95%.

Remark: Solutions in reagent-grade acetone; Water quality parameters

monitored throughout test.

Result: 96 hr C.I. = 0.12 - 0.16 mg/l;

24 hr LC50 = 0.28 mg/l; 48 hr LC50 = 0.18 mg/l

Test condition: carrier-acetone; 15L water; 10 fish/vessel; length =

3.7 cm; no food; no aeration; temp = 12C

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (19)

Type: static

Species: Lepomis macrochirus (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: yes

LC50: .4

Method: other: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians.

Year: 1977 GLP: no data Test substance: other TS: Santoflex 13 Lot# KD03-017, purity: >95%.

Remark: Solutions in reagent-grade acetone; Water quality parameters

monitored throughout test.

Result: 96 hr C.I. = 0.32 - 0.5 mg/l;

24 hr LC50 = 0.65 mg/l;48 hr LC50 = 0.45 mg/l

Test condition: carrier-acetone; 15L water; 10 fish/vessel; length =

3.8 cm; no food; no aeration; temp = 22C

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (19)

- 15/57 -

Type: static

Species: Brachydanio rerio (Fish, fresh water)

Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: no

LC0: 5 LC100: 100

Method: other: see remarks

Year: 1984 GLP: no

Test substance: other TS: technical grade 6PPD

Remark: following OECD 203

The powdered test substance was dispersed in water. LC-values given above are nominal concentrations: weight of

the dispersed substance per liter water.

Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

20-NOV-2001 (20)

Type: flow through

Species: Pimephales promelas (Fish, fresh water)

Exposure period: 28 day

Unit: mg/l Analytical monitoring: yes

LC50: = .15

Method: other: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians.

Year: 1984 GLP: yes

Test substance: other TS: Santoflex 13 purity: >95%.

Remark: C.I. = 0.13 - 0.17 mg/l; 48 hr LC50 = 2 mg/l; 6, 7 and 8 day LC50 = 0.35 mg/l; 19, 20, 21 day LC50 = 0.17 mg/l

Tests in well water; Stock solutions in acetone; Water quality

parameters monitored throughout test.

parameters monitored throughout tes

Result: 28D C.I. = 0.13 - 0.17 mg/l;

48 hr LC50 = 2 mg/l;

6, 7 and 8 day LC50 = 0.35 mg/l; 19, 20, 21 day LC50 = 0.17 mg/l (1) valid without restriction

Reliability: (1) valid without restriction

GLP guideline study

20-NOV-2001 (21)

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#### 4.2 Acute Toxicity to Aquatic Invertebrates

Type: static

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: yes

NOEC: = .56 EC50: = .82

Method: other: EPA Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates and Amphibians

Year: GLP: yes

Test substance: other TS: Santoflex 13, purity: >95%

Remark: Solutions in reagent-grade acetone; Water quality parameters

monitored throughout test.

Result: C.I. for 48 hr EC50=0.71-0.94 mg/l;

24 hr EC50=1 mg/l

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (22)

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = .4 EC50: = .79

Method: OECD Guide-line 202, part 1 "Daphnia sp., Acute

Immobilisation Test"

Year: 1984 GLP: no data

Test substance:

Remark: C.I. for EC50 = 0.7 - 0.91 mg/l; 24 hr EC50=1.6 mg/l;

48 hr EC50=0.79 mg/l; in presence of food 48 hr EC50=

1.3 mg/l and NOEC=0.4 mg/l

Source: MonsantoBayer AG Leverkusen
Test condition: carrier-acetone; no food
Reliability: (1) walid without restriction

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (23)

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Date: 20-NOV-2001
4. Ecotoxicity ID: 793-24-8

Type:

Species: Daphnia magna (Crustacea)

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = .25 EC50: = .51

Method: OECD Guide-line 202, part 1 "Daphnia sp., Acute

Immobilisation Test"

Year: 1984 GLP: no data

Test substance:

Remark: the test solution was allowed to age 40 hours before test

48 hr EC50>1 mg/l and NOEC>1 mg/l

Source: MonsantoBayer AG Leverkusen Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (24)

Type:

Species: other: Chironomus tentans

Exposure period: 48 hour(s)

Unit: mg/l Analytical monitoring: no

NOEC: = .6 EC50: = .99

Method: other: EPA. Methods for Acute Toxicity Tests with Fish,

Macroinvertebrates, and Amphibians. EPA-660/3-75-009.

Year: 1975 GLP: no data

Test substance:

Remark: C.I. for EC50=0.6-1.25 mg/l; 24hr EC50=1.25 mg/l

Source: MonsantoBayer AG Leverkusen

Test condition: water solubility was exceeded at three highest concen-

trations; larvae 10-14 days old; room temp

30-MAY-1994 (25)

4.3 Toxicity to Aquatic Plants e.g. Algae

Species: Selenastrum capricornutum (Algae)

Endpoint: biomass
Exposure period: 96 hour(s)

Unit: mg/l Analytical monitoring: yes

EC50: = .6

Method: other: EPA Selenastrum capricornutum Algal Assay Test Year: GLP: no data

Test substance: other TS: Santoflex 13 (Monsanto) purity: >95%

Remark: Phytotoxicity maxed at 48 hours; test solutions in acetone

Result: 96 hr C.I. 0.2-2 mg/l;

in vivo chlorophyll results-

24hr EC50=2.0 mg/l, 48hr EC50=0.5 mg/l, 72hr EC50=0.5 mg/l, 96hr EC50=0.6 mg/l

Test condition: temp=24C; 4000 lux; Algal Assay media; "cool" white lights;

init. inoc.=10000 cells/ml

Reliability: (1) valid without restriction

Guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (26) (27)

4.4 Toxicity to Microorganisms e.g. Bacteria

Type:

Species: activated sludge

Exposure period: 3 hour(s)

Unit: mg/l Analytical monitoring:

EC50: 420

Method: ISO 8192 "Test for inhibition of oxygen consumption by

activated sludge"

Year: GLP: no

Test substance: other TS

Source: Bayer AG Leverkusen Test substance: technical grade 6PPD

01-DEC-1992 (20)

# 4.5 Chronic Toxicity to Aquatic Organisms

# 4.5.1 Chronic Toxicity to Fish

Species: Endpoint:

Exposure period:

Analytical monitoring: Unit:

Method:

GLP: Year:

Test substance:

Remark: no information

Source: Bayer AG Leverkusen

06-FEB-1992

## 4.5.2 Chronic Toxicity to Aquatic Invertebrates

Species: Endpoint:

Exposure period:

Analytical monitoring: Unit:

Method:

Year: GLP:

Test substance:

Remark: no information

Source: Bayer AG Leverkusen

06-FEB-1992

## TERRESTRIAL ORGANISMS

#### 4.6.1 Toxicity to Soil Dwelling Organisms

Type: Species: Endpoint:

Exposure period:

Unit: Method:

Year: GLP:

Test substance:

Remark: no information

Source: Bayer AG Leverkusen

06-FEB-1992

# 4.6.2 Toxicity to Terrestrial Plants

Species: Endpoint:

Expos. period:

Unit: Method:

Year: GLP:

Test substance:

Remark: no information
Source: Bayer AG Leverkusen

06-FEB-1992

# 4.6.3 Toxicity to other Non-Mamm. Terrestrial Species

# 4.7 Biological Effects Monitoring

Remark: no information

Source: Bayer AG Leverkusen

06-FEB-1992

# 4.8 Biotransformation and Kinetics

Type:

Remark: no information

Bayer AG Leverkusen Source:

06-FEB-1992

# 4.9 Additional Remarks

### 5.1 Acute Toxicity

### 5.1.1 Acute Oral Toxicity

Type: LD50 Species: rat

Strain: Sprague-Dawley Sex: male/female

Number of

Animals: 10

Vehicle:

Value: > 5000 mg/kg bw

Method: other: EPA/TSCA Acute Oral Toxicity and the EEC Methods for

Determining Toxicity, Part B.1, No. L 251/96 Sept. 1984

Year: GLP: yes

Test substance: other TS: 6PPD Ref# 4065459 solid, purity: 97.6%

Remark: Following a range-finding study, 6PPD was fed to a group of

five male and five female rats in a single oral dose of 5000

mg/kg body weight. Rats were observed daily and weighed

weekly. 2 males and I female died prior to sacrifice. A gross necropsy examination was performed on all surviving animals at sacrifice on Day 15. Clinical findings included decreased fecal output, fecal/urine stains, rough coat, piloerection and

soft stools. One male and three females showed weight loss; all other animals gained weight. Most notable internal necropsy finding was black, hard material in the stomach contents. Findings in animals that died included discolored mucoid contents throughout the digestive system with reddened

mucosa/dark red foci of the stomach.

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (28)

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 3340 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: undiluted Source: Bayer AG Leverkusen

08-DEC-1992 (29)

-

Type: LD50 Species: rat

Strain:
Sex:
Number of
 Animals:
Vehicle:

Value: = 2500 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (30)

Type: LD50 Species: rat

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 3580 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: purity 95.7 % Source: Bayer AG Leverkusen

08-DEC-1992 (31)

Type: LD50 Species: mouse

Strain:
Sex:
Number of
Animals:
Vehicle:

Value: = 3200 mg/kg bw

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (30)

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Date: 20-NOV-2001 5. Toxicity ID: 793-24-8

Type: LD50

Species: Strain: Sex: Number of Animals: Vehicle:

Value: = 1120 mg/kg bw

Method:

GLP: Year:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (32)

5.1.2 Acute Inhalation Toxicity

5.1.3 Acute Dermal Toxicity

Type: LD50 Species: rabbit

Strain: New Zealand white

Sex: male/female

Number of Animals:

Vehicle: other: undiluted Value: > 7940 mg/kg bw

other: Defined Lethal Dose Method:

Year: GLP: no data

Test substance: other TS: CP 22423 Lot# KC07-298, purity: >95%.

Remark: The undiluted test article was applied to the shaved skin of male and female rabbits at dose levels ranging from 3160 to 7940 mg/kg/bw. Clinical signs were reduced appetite and activity for three to seven days. All animals survived. Autopsy results showed that all viscera appeared normal.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flaq: Critical study for SIDS endpoint

20-NOV-2001 (31)

LDLo Type: Species: rabbit

Strain: Sex: Number of

Animals: Vehicle:

Value: 3160 - 5010 mg/kg bw

Method:

GLP: Year:

Test substance: other TS: undiluted Bayer AG Leverkusen

08-DEC-1992 (29)

5.1.4 Acute Toxicity, other Routes

-

5.2 Corrosiveness and Irritation

5.2.1 Skin Irritation

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: slightly irritating

EC classificat.:

Method: Draize Test

Year: GLP:

Test substance: other TS: undiluted

Remark: method: the data were scored according to the method of

Draize et al. (1944), 24 h exposure, then skin rinsed with

warm water and soap, observation period 5 days

Source: Bayer AG Leverkusen

08-DEC-1992 (29)

Species: rabbit

Concentration:

Exposure:
Exposure Time:
Number of
Animals:
PDII:

Result: slightly irritating

EC classificat.:

Method: Draize Test

Year: GLP:

Test substance: other TS: 12.5 and 125 mg 6PPD dispersed in 0.5 g vaseline

(2.5 and 25 %)

Remark: method: after 24 h and 72 h examination

Source: Bayer AG Leverkusen

08-DEC-1992 (33)

- 25/57 -

Date: 20-NOV-2001 5. Toxicity ID: 793-24-8

Species: rabbit

Concentration:

Exposure: Exposure Time: Number of Animals: PDII:

Result: moderately irritating

EC classificat.:

Method: Draize Test

Year: GLP:

Test substance: other TS: 25 mg 6PPD dispersed in 0.5 ml olive oil

Remark: method: after 24 h and 72 h examination

Source: Bayer AG Leverkusen

08-DEC-1992 (33)

Species: rabbit

Concentration:

Exposure: Exposure Time: Number of Animals: PDII:

Result:

not irritating

EC classificat.:

Method: other: (see remarks)

Year: GLP:

Test substance:

method: 0.5 ml, semi-occlusive, clipped intact and Remark: abraded skin, 24 h exposure, observation period 7 days, scoring in accordance with the Federal Hazardous Substance

Act, 21 CFR, paragraph 191.11 (1964)

Source: Bayer AG Leverkusen

08-DEC-1992 (31)

5.2.2 Eye Irritation

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment: Number of Animals:

Result: slightly irritating

EC classificat.:

Method: other: (see remarks)

Year: GLP:

Test substance: other TS: undiluted

method: 0.1 ml in the conjunctival sac of the right eye of Remark:

each of 3 rabbits, 24 h exposure, then eyes rinsed with warm isotonic saline solution, observation period 5 days, the

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Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

data were scored according to the method of Draize et al.

(1944)

Source: Bayer AG Leverkusen

08-DEC-1992 (29)

Species: rabbit

Concentration:

Dose:

Exposure Time: Comment: Number of Animals:

Result: slightly irritating

EC classificat.:

Method: other: (see remarks)

Year: GLP:

Test substance:

Remark: method: 0.1 ml in the conjunctival sac, observation

period 7 days, scoring in accordance with the Federal Hazardous Substance Act, 21 CFR, paragraph 191.12 (1964)

Source: Bayer AG Leverkusen

08-DEC-1992 (31)

5.3 Sensitization

Type: Guinea pig maximization test

Species: guinea pig

Number of Animals: Vehicle:

Result: sensitizing

Classification:

Method:

Year: GLP:

Test substance: other TS: 6PPD in olive oil or vaseline
Remark: 50 % sensitization (challenge with 0.05 %),
90 % sensitization (challenge with 0.5 %)

Source: Bayer AG Leverkusen

08-DEC-1992 (33)

Type: Patch-Test Species: human

Concentration: Induction 50 %

Number of

Animals: 50

Vehicle: Result:

Classification:

Method: other: Modified Draize

Year: GLP: Test substance: other TS: PPD; purity not stated

Remark: PPD was patch tested on 50 human volunteers at a concentration

of 50% w/v in dimethylphthalate. 5 of the 50 subjects showed skin reactions during the 3-week induction phase of the study.

5 of 50 subjects showed skin reactions in the challenge phase.

20-NOV-2001 (34)

Type: Patch-Test

Species: human

Number of Animals: Vehicle: Result:

Classification:

Method: other: Repeated Insult Patch Test Year: GLP:

Test substance: other TS: a 0.1 % W/V solution in dimethylphthalate

Remark: 0/50 volunteers had a positive test result

Source: Bayer AG Leverkusen

08-DEC-1992 (35)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS

Remark: 0/50 (for each rubber sample) human subjects not previously

exposed to test rubber formulations had a positive patch

test result

Source: Bayer AG Leverkusen

Test substance: 2 parts 6-PPD per hundred parts rubber, unvulcanized

2 parts 6-PPD per hundred parts rubber, vulcanized

20-MAY-1992 (36)

Type: Patch-Test
Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 6PPD as additive

Remark: 17/50 subjects showed a positive reaction after challenge

Source: Bayer AG Leverkusen

08-DEC-1992 (37)

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Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 2 parts 6PPD per hundred parts

rubber

Remark: 2/4 volunteer subjects who had reacted to previous rubber

samples, had a positive patch test result

Source: Bayer AG Leverkusen

08-DEC-1992 (38)

Type: Patch-Test Species: human

Number of Animals: Vehicle: Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 2 parts 6PPD per hundred parts

rubber

Remark: 5/10 volunteer subjects who had reacted to previous rubber

samples, had a positive patch test result

Source: Bayer AG Leverkusen

08-DEC-1992 (39)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 6PPD as additive

Remark: 3/10 volunteer subjects, all of whom had been previously

sensitized to a rubber sample, had a positive patch test

result

Source: Bayer AG Leverkusen

08-DEC-1992 (40)

- 29/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: samples with 1, 2 and 3 parts 6PPD per hundred parts

rubber

Remark: 9/10 (for each rubber sample) volunteer subject who had

reacted to previous rubber samples, had a positive patch

test result

Source: Bayer AG Leverkusen

08-DEC-1992 (41)

Type: Patch-Test

Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 6PPD as additive

Remark: 4/50 subjects showed a positive reaction after challenge

Source: Bayer AG Leverkusen

08-DEC-1992 (42)

Type: Patch-Test

Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: a rubber sample with 2 parts 6PPD per hundred parts

rubber

Remark: 0/50 volunteer subjects, not previously associated

with either chemical had a positive patch test result

Source: Bayer AG Leverkusen

08-DEC-1992 (43)

- 30/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

5. TOXICIES

Type: Patch-Test Species: human

Number of Animals: Vehicle: Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: 1 % Santoflex 13 in petrolatum

Remark: No skin reactions were noted in a 6-week study on 94 human volunteers. The induction phase consisted of the application

of 1% 6PPD in petrolatum to the same site, 3x/week for three weeks. In the challenge phase, the test article was applied

at a previously unpatched site.

20-NOV-2001 (44)

Type: Patch-Test Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: 50 % w/v Santoflex 13 in dimethylphthalate Remark: 50 human volunteers were patch tested with 50 % w/v

Santoflex 13 in dimethylphthalate; five of the 50 individuals showed reactions in the 3-week induction phase and 5 of 50 showed reactions in the challenge

phase

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (45)

Type: Patch-Test
Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: no data

Remark: 6/9 contact dermatitis patients showed a positive reaction

with 6PPD

Source: Bayer AG Leverkusen

17-AUG-1998 (46)

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Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

5. Tokiete, 12. 755 21 0

Type: Patch-Test

Species: human Number of

Animals: Vehicle: Result:

Classification:

Method:

Year: GLP:

Test substance: no data

Remark: 6/135 contact dermatitis patients showed a positive reaction

with 6PPD

Source: Bayer AG Leverkusen

17-AUG-1998 (47)

Type: no data Species: human

Number of
Animals:
Vehicle:
Result:

Classification:

Method:

Year: GLP:

Test substance: other TS: 2 % in lanolin

Remark: 15/15 IPPD-allergic patients were positive in the test with

6-PPD

Source: Bayer AG Leverkusen

08-DEC-1992 (33)

Type: other: (see remarks)

Species: guinea pig

Number of
Animals:
Vehicle:

Result: not sensitizing

Classification:

Method: other: (see remarks)

Year: GLP:

Test substance:

Remark: method: application daily for 20 days (50 % paste), back,

for the challenge different concentrations 10, 20, 30, 50 and 100 %) were applied to new areas of the back (no further

data available)

Source: Bayer AG Leverkusen

08-DEC-1992 (30)

-

# 5.4 Repeated Dose Toxicity

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: oral feed Exposure period: 13 w

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 250, 1000 or 2500 ppm (19, 75 or 188 mg/kg b.w./d)

Control Group: yes, concurrent no treatment

NOAEL: 250 ppm

Method: other: EHL Protocol 85087 Ref: Multiple Comparison Procedure

for Comparing Several Treatments with a Control (1955)

Year: GLP: yes

Test substance: other TS: Santoflex 13 Lot#KE06-121, purity: 97.1%

Result: Santoflex 13 was administered in feed to groups of

Santoflex 13 was administered in feed to groups of 6 week old male and female rats at the above levels. Analyses via GC verified feeding levels of 0, 230, 950 and 2300 ppm. All animals survived the length of the study. Signs of toxicity during the study were limited to reduced feed consumption/body weight gain in the high-dose males and females and mid-level males. Anemia, lymphocytopenia and thrombocytosis were present in males and females, primarily at the two highest dose levels. Increases in total bilirubin in males, and total protein, albumin, globulin, calcium and/or cholesterol in both sexes were noted in high and some mid-dose level animals. Increased liver weights were observed at the two highest dose levels. There were no gross or microscopic lesions attributed to consumption of the test material. Females at low dose levels exhibited mild anemia at the interim sampling period

levels exhibited mild anemia at the interim sampling period, but all recovered by the end of the study. Therefore, the

NOEL was considered to be 250 ppm.

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (48) (49)

- 33/57 -

Date: 20-NOV-2001 5. Toxicity ID: 793-24-8

Species: rat Sex: male/female

Strain: Sprague-Dawley Route of admin.: inhalation

Exposure period: 4 w (20 exposures)

Frequency of

6 h/d treatment:

Post. obs.

period: no data

0.054, 0.236 or 0.477 mg/l Doses: Control Group: yes, concurrent no treatment

Method: other: Subacute Dust Inhalation Study IBT #8562-09721

(Audited)

Year: GLP: yes

Test substance: other TS: Santoflex 13 Powder Lot #KD03-017, purity: 97.1% Result: 4 groups of 5 male and 5 female young adult albino rats were

exposed to either zero, low, intermediate or high dust

concentrations of the test article. Test dusts were suspended in streams of clean, dry air, and introduced through the top center of exposure chambers and exhausted out the bottom. GC analytical testing confirmed concentrations and total weight of test dusts. All but one animal survived until sacrifice on Day 28. Hypoactivity was noted in all test groups. Mid and high-dose animals exhibited swollen snouts and scratching. Mean body weights of treated animals compared favorably with those of controls. Results of gross necropsy indicated increased liver and kidney weights of treated animals over those of controls. Lung weights were reduced in high-dose makes and mid-dose females. Mid-dose treated males exhibited increased spleen weights. No significant differences were noted in the weights of the brains, gonads and hearts of treated animals when compared to controls. No gross or histopathologic alterations attributed to the test article

were observed in any of the treated animals.

Mean corpuscular hemoglobin was reduced in high-dose males; elevations in SGPT and lowered glucose levels in mid- and high-dose males were correlated with increased relative liver weights; no treatment rela-

ted gross lesions were noted at necropsy.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Critical study for SIDS endpoint Flag:

20-NOV-2001 (50)

Species: rat Sex: male/female

Strain: other: Charles River CD

Route of admin.: oral feed Exposure period: 24 months

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 100, 300, 1000 ppm (8, 23, 75 mg/kg b.w./day)

Control Group: yes, concurrent no treatment

NOAEL: 23 mg/kg LOAEL: 75 mg/kg

Method: other: 2-Year Chronic Oral Toxicity IBT Protocol # 622-05400A

(1974)

Year: GLP: yes Test substance: other TS: 6PPD. Powder, purity: 96.9%

Remark: hematology, clinical chemistry and urinalysis were conducted

at 3, 6, 12 and 24 months, the calculation of the dose levels is based on 1 ppm corresponds to 0.075 mg/kg b.w.; 50

male and 50 female rats per group.

Result: 6PPD was fed at the above doses to groups of 200 male and 200

female rats over a two-year period, beginning when the males were 28 days old and the females 29 days old. Dose levels were

verified by GC analysis. Body weight, food consumption, behavior, hematology, blood chemistry and urinalysis results were recorded throughout the study. Complete gross necropsies were conducted on all animals found dead, on all animals sacrificed in extremis, and on all remaining animals at 24

months.

All organs or tissues with grossly visible lesions were submitted for histologic examination. Statistical reductions in body weight were noted in high-dose males during Weeks 1-5. High-dose females exhibited statistically reduced body weights throughout the study. Body weights and weight gain of the midto low-dose animals compared favorably to controls. Frequency and distribution of deaths during the study were similar

to low-dose animals compared favorably to controls. Frequency and distribution of deaths during the study were similar between treated animals and controls. Gross pathological examination of animals that died during the study did not reveal any relation to death and the test article. There were no unusual behaviors noted in test animals during the study. A significant reduction in erythrocyte counts was noted in high-dose males at 3 months and in high-dose females at 3, 6, and 9 months. However, the same animals had erythrocyte counts similar to controls at all subsequent blood

collections. Hemoglobin concentration, while still considered to be within normal range, was statistically reduced for

high-dose males at 3, 12 and 18 months. High-dose females exhibited similar reductions at 6, 12 and 18 months.

Hematocrit values among high-dose animals were significantly lower than controls, and were at the lower limits at 3 and 12 months for males, and 3,6 and 12 months for females.

Hematocrit values in these animals exhibited a slight increase at 18 and 24 months. Urinalysis studies, which included monitoring of glucose, albumin, microscopic elements, pH and specific gravity, were similar for both treated and control

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groups throughout the study. Gross pathological examination of animals sacrificed at 24 months revealed similar findings for both treated and control groups. Statistical analysis of absolute organ weights, organ to body weight ratios and organ to brain weight ratios compared favorably across the test and control groups, and were within the range of expected values for albino rats of this age and strain. Histopathological examination of organs and tissue taken from high-dose animals and controls at 24 months revealed no treatment-related lesions. Any lesions noted were from those of naturally-occurring diseases, and were noted in both populations. Microscopic examination of suspect lesions from all sacrificed animals and also those that died during the study. No differences were noted between test and control rats

as to the organ system involved, type or classification of neoplasms..

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (51) (52)

Species: rat Sex: male/female

Strain: no data Route of admin.: oral feed

Exposure period: after 12 months interim sacrifice (no further data)

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 50, 250 or 1500 ppm (4, 20 or 120 mg/kg bw/d)

Control Group: yes

Method:

Year: GLP: no data

Test substance: other TS: Santoflex 13

Remark: The NOEL for chronic toxicity was determined to be

50 ppm, and a NOEL for oncogenic effects was deter-

mined to be at least 1500 ppm

Result: decreased body weights in mid- and high-exposure fe-

males and high-exposure males; various hematological changes in mid- and high-exposure females and highexposure males; some high-exposure male and female serum chemistry alterations (increased cholesterol, total protein, globulin and calcium); absolute and relative liver weights were increased for mid-exposure male rats at study termination and for high-exposure male and female rats after one year of exposure and at the end of the study; histopathological examination revealed pigment in the hepatocytes and reticuloendothelial cells of high-exposure females; mean absolute and relative kidney weights were also statistically significantly increased for high-exposure males and females compared to controls at the 12-month interim sacrifice only; a slight increase in the severity but not the incidence of chronic nephropathy was noted for high-expo- 36/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

sure males and females compared to controls at both interim and terminal sacrifice periods; high exposure males demonstrated increased absolute and relative spleen

weights compared to controls at the 12-month exposure period only; neoplastic findings were similar between

control and Santoflex 13-treated animals

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (53)

Species: rat Sex: no data

Strain:

Route of admin.: gavage Exposure period: 24 days

Frequency of

treatment: once a day

Post. obs.

period: no data

Doses: 250 mg/kg b.w./day for the first 4 days, thereafter being

increased 50 % every 5 days, no further data available

Control Group: yes

Method:

Year: GLP:

Test substance:

Result: no death, body weight gain within the normal range,

increased oxygen consumption, suppression of the central nervous system and of the synthezising function of the liver

(content of hippuric acid in a 24 h urine sample was decreased), decreased ascorbic acid content in the liver

Source: Bayer AG Leverkusen

08-DEC-1992 (30)

### 5.5 Genetic Toxicity 'in Vitro'

Type: Ames test

System of

testing: Salmonella typhimurium TA-1535 TA-1537 TA-1538 TA-98 TA-100

Concentration: 0.001, 0.01, 0.1, 1.0 and 5.0 micrograms/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Ames Plate Test (Overlay method) 1975; OECD 471

equivalent

Year: GLP: yes
Test substance: other TS: 6PPD #BIO76-277, purity: >96%

Remark: Stock solutions prepared in DMSO. No evidence of mutagenic

activity in any assay conducted with or without activation using the S-9 homogenate from Arochlor-induced rat livers.

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (54)

-

Type: Ames test

System of

testing: Salmonella typhimurium TA1535, TA1537, TA1538, TA98, TA100 Concentration: 0.167, 0.500, 1.67, 5.00, 16.7 and 50.0 micrograms/plate

Cytotoxic Conc.: Precipitation conc: >500 micrograms/plate

Metabolic

activation:

Result:

Method: other: Revised Method for the Salmonella Mutagenicity Test

(1983), Maron, D.M. and Ames, B.N.

Year: GLP: yes

Test substance: other TS: 6PPD purple solid #4065461, purity: >96%

Remark: Stock solutions prepared in DMSO. All tester strains contained

a uvrB deletion mutation and an rfa mutation. Cytotoxicity of test article was determined in a screening test on duplicate cultures of TA1538 and TA100 in the absence of S9. In the

definitive assay, inhibited growth was observed at

concentrations >5.00, both with and without S9 activation. Revertant frequencies for all doses, in all strains, both with and without metabolic activation were equal to or less than

those of controls. Results for the test article were

negative under the test conditions.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (55)

Type: Gene mutation in Saccharomyces cerevisiae

System of

testing: Saccharomyces cerevisiae D4

Concentration: 0.001, 0.01, 0.1, 1.0 and 5.0 micrograms/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Ames Plate Test (Overlay method) 1975; OECD 471

equivalent

Year: GLP: yes

Test substance: other TS: 6PPD #BIO76-277, purity: >96% Remark: Stock solutions prepared in DMSO. No evidence

mark: Stock solutions prepared in DMSO. No evidence of mutagenic activity in any assay conducted with or without activation using the S-9 homogenate from Arochlor-induced rat livers.

Reliability: (1) valid without restriction

GLP guideline study

Flag: Critical study for SIDS endpoint

20-NOV-2001 (54)

- 38/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Mammalian cell gene mutation assay

System of

testing: Mouse lymphoma cells (L5178Y TK+/-)

Concentration: 0.25, 0.5, 1.0, 2.0, 4.0 or 8.0 micrograms/ml

Cytotoxic Conc.: With metabolic activation: 33 micrograms/ml; Without

metabolic activation: > 4 micrograms/ml

Metabolic

activation: with and without

Result: negative

Method: other: OECD 476 equivalent

Year: GLP: yes Test substance: other TS: 6PPD/CP22423 , purity: >96%

Remark: Negative for ability to induce forward mutations at the TK

locus.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (56)

Type: Mammalian cell gene mutation assay

System of

testing: Chinese hamster ovary cells (CHO/HGPRT)

Concentration: up to 5 ug/ml without S9-mix, up to 15 ug/ml with S9-mix Cytotoxic Conc.: With metabolic activation: 9 micrograms/ml; Without

metabolic activation: 4 micrograms/ml; Solubility limit of

test article = 333 micrograms/ml

Metabolic

activation: with and without

Result: negative

Method: other: CHO/HGPRT Mutation Assay (1979) Hsie, et.al.

Year: GLP: yes

Test substance: other TS: 6PPD purple pellets lot# KH04, purity: 96%

Remark: 6PPD was tested in CHO cells at different S9 concentrations up

up to cytotoxic concentrations in two range-finding, one initial and one confirmatory experiments. The cytoxicity of the test article decreased with increasing S9 concentrations.

No statistically significant mutagenicity was observed. 6PPD

is not considered mutagenic to CHO cells under test

conditions.

Reliability: (1) valid without restriction Flag: Critical study for SIDS endpoint

20-NOV-2001 (57)

-

Type: Unscheduled DNA synthesis

System of

testing: primary rat hepatocytes

Concentration: 0.1, 0.5, 1, 5, 10, 50, 100, 500, 1000 and 5000 micrograms/ml

Cytotoxic Conc.: 50 micrograms/ml

Metabolic

activation: without Result: negative

Method: other: Williams, G.M., 1977. Detection of Chemical Carcinogens

by Unscheduled DNA Synthesis in Rat Liver Primary Cell

Cultures

Year: GLP: yes

Test substance:

Remark:

other TS: 6PPD purple pastilles Lot# KH04-70, purity: 96% Reagent grade Acetone (1%) as solvent. 6PPD was examined for genotoxicity in the UDS Assay. Primary rat lever cell cultures used for both the preliminary and replicate experiments were derived from the livers of two adult male Fischer-344 rats (13)

and 18 weeks old, respectively).

Quantitative autoradiographic grain-counting was performed using an ARTEK Model 980 colony counter interfaced with a Zeiss Universal Microscope via an ARTEK TV camera. Data were fed directly to a VAX computer. Cytotoxicity was observed at concentrations of 50 micrograms/ml and above in both the preliminary and replicate experiments. UDS was measured at concentrations of the test article between 0.1 and 10

concentrations of the test article between 0.1 and 10 micrograms/ml in both experiments. The net grain counts were negative at each concentration of the test compound, in the solvent control, and in the medium control, in contrast to the strong positive response produced in both experiments by the positive control. These results indicate that 6PPD is not a

genotoxic agent under the conditions of the in vitro rat

hepatocyte DNA repair assay.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (58)

Type: Cytogenetic assay

System of

testing: Chinese hamster ovary cells (CHO)

Concentration:
Cytotoxic Conc.:

Metabolic

activation: no data Result: negative

Method: other: chromosomal aberrations
Year: GLP:

Test substance:

Remark: no further data available Source: Bayer AG Leverkusen

20-NOV-2001 (59)

- 40/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

.. Tokietey 110. 773 21 0

Type: Cytogenetic assay

System of

testing: Chinese hamster ovary cells

Concentration: up to 15 ug/ml

Cytotoxic Conc.:

Metabolic

activation: no data

Result: Method:

Year: GLP: no data

Test substance: other TS: Santoflex 13

Remark: effects: Santoflex 13 showed a marginal potential for

inducing chromosomal aberrations
type: chromosomal aberration assay

Source: MonsantoBayer AG Leverkusen

20-NOV-2001 (60)

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration: up to 1000 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method: other: Ames Salmonella/Microsome (EPA/OECD)

Year: 1984 GLP:

Test substance: other TS: Flexzone 7F Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions

21-OCT-1999 (61)

Type: Ames test

 ${\tt System} \ {\tt of} \\$ 

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537, TA

1538

Concentration:
Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

Remark: no further data available Source: Bayer AG Leverkusen

08-DEC-1992 (62)

- 41/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Ames test

System of

testing: Salmonella typhimurium TA 98, TA 100, TA 1535, TA 1537

Concentration: up to 200 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (63)

Type: Ames test

System of

testing: Salmonella typhimurium

Concentration:
Cytotoxic Conc.:

Metabolic activation:

Dogult:

Result: negative

Method:

Year: GLP:

Test substance:

Remark: no further data available Source: Bayer AG Leverkusen

08-DEC-1992 (64) (65) (66)

Type: Ames test

System of

testing: Salmonella typhimurium (no further data)

Concentration: up to 500 ug/plate

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP: no data

Test substance: other TS: Santoflex 13

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (67)

- 42/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Mammalian cell gene mutation assay

System of

testing: Chinese hamster ovary cells (CHO/HGPRT)

Concentration: up to 0.6 ug/ml without S-9 mix, up to 55 ug/ml with S-9 mix

Cytotoxic Conc.:

Metabolic

activation: with and without

Result: negative

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (68)

Type: Mitotic recombination in Saccharomyces cerevisiae

System of

testing: Saccharomyces cerevisiae D4

Concentration: no data

Cytotoxic Conc.:

Metabolic

activation: no data Result: negative

Method:

Year: GLP: no data

Test substance: other TS: Santoflex 13

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (69) (70)

Type: Sister chromatid exchange assay

System of

testing: Chinese hamster ovary cells (CHO)

Concentration:
Cytotoxic Conc.:

Metabolic

activation: no data Result: negative

Method:

Year: GLP:

Test substance:

Remark: no further data available Source: Bayer AG Leverkusen

08-DEC-1992 (59)

- 43/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Type: Unscheduled DNA synthesis

System of

testing: primary rat hepatocyte Concentration: up to 1000 ug/well

Cytotoxic Conc.:

Metabolic

activation: without Result: negative

Method:

Year: GLP:

Test substance:

Source: Bayer AG Leverkusen

08-DEC-1992 (71)

Type: Unscheduled DNA synthesis

System of

testing: primary rat hepatocytes

Concentration: up to 1000 ug/ml

Cytotoxic Conc.:

Metabolic

activation:

Result: negative

Method:

Year: GLP: no data

Test substance: other TS: Flexzone 7F

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (72)

5.6 Genetic Toxicity 'in Vivo'

Type: Cytogenetic assay

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage

Exposure period: 6, 18 and 30 hours Doses: 1000 mg/kg bw

Result: negative

Method: other: EPA Health Effects Test Guidelines EPA 560/6-82-09

Year: 1984 GLP: yes
Test substance: other TS: 6PPD Lot# KJ09-165, purity: 96%

Remark: Not clastogenic under test conditions. Mild to severe

pharmacotoxic effects observed in test animals indicated that the test article was administered near the maximum tolerated

dose.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (73)

- 44/57 -

Date: 20-NOV-2001 5. Toxicity ID: 793-24-8

Type: Cytogenetic assay

Species: Sex: male mouse

Strain:

Route of admin.: i.p.

Exposure period: twice within 24 hours Doses: 100 and 200 mg/kg bw Result: negative

other: no data Method:

GLP: no data Year:

Test substance: no data

Result: no induction of chromosomal abnormalities

Source: Bayer AG Leverkusen

(74)20-NOV-2001

Type: Micronucleus assay

Species: mouse Sex: male/female

Strain: CD-1 Route of admin.: i.p. Exposure period: 1 day Doses: 1000 mg/kg Result: negative

Method:

Year: GLP:

Test substance:

Remark: clinical signs were assessed

Source: no increased number of micronucleated erythrocytes

Bayer AG Leverkusen

20-NOV-2001 (75)(76)

Type: Micronucleus assay

Species: mouse Sex: male

Strain:

Route of admin.: i.p.

Exposure period: twice within 24 hours 100, 150 and 200 mg/kg bw Doses:

negative Result: other: no data Method:

Year: GLP: no data

Test substance: no data

Result: no induction of micronucleated erythrocytes in bone marrow

Bayer AG Leverkusen Source:

20-NOV-2001 (74) - 45/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

5.7 Carcinogenicity

Species: rat Sex: male/female

Strain: other: Charles river CD

Route of admin.: oral feed Exposure period: 24 months

Frequency of

treatment: daily

Post. obs.

period: no

Doses: 100, 300, 1000 ppm (8, 23, 75 mg/kg b.w./day)

Result:

Control Group: yes, concurrent no treatment

Method:

Year: GLP:

Test substance:

Remark: the calculation of the dose levels is based on 1 ppm

corresponds to 0.075 mg/kg b.w.; 50 male and female rats

per group

Result: the number and type of neoplastic and nonneoplastic lesions

were comparable between groups

Source: Bayer AG Leverkusen

08-DEC-1992 (77)

Species: rat Sex: male/female

Strain: no data Route of admin.: oral feed

Exposure period: after 12 months interim sacrifice (no further data)

Frequency of

treatment: daily

Post. obs.

period: no data

Doses: 50, 250 or 1500 ppm (4, 20 or 120 mg/kg bw/d)

Result:

Control Group: yes

Method:

Year: GLP: no data

Test substance: other TS: Santoflex 13

Remark: a NOEL for oncogenic effects was determined to be at

least 1500 ppm

Result: neoplastic findings were similar between control and

Santoflex 13-treated animals (no further data)

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (53)

- 46/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Species: other: (see remarks) Sex:

Strain:

Route of admin.:
Exposure period:
Frequency of
 treatment:
Post. obs.
 period:
Doses:

Control Group:

Method:

Result:

Year: GLP: yes

Test substance:

Remark: BALB/3T3 cells; cell transformation assay under

nonactivation conditions

Result: negative

Source: Bayer AG Leverkusen

08-DEC-1992 (78)

5.8 Toxicity to Reproduction

Type: Fertility

Species: rat Sex: male/female

Strain: Sprague-Dawley

Route of admin.: gavage

Exposure Period: Males: 42 or 49 days, Females: 14 days prior to mating

through Day 7 of gestation

Frequency of

treatment: daily
Premating Exposure Period
male: 28 days.
female: 14 days

Duration of test:

Doses: 0, 40, 200 or 1000 ppm Control Group: yes, concurrent vehicle

NOAEL Parental: > 1000 ppm NOAEL F1 Offspr.: > 1000 ppm

Method: other: Fertility Study and Early Embrionic Development to

Implantation in Rats, DRL

Year: 1998 GLP: no data

Test substance: other TS: CD-13, purity >98%

Remark: The test article is being evaluated as a new diagnostic drug

of Helicobacter pylori. To this end, several reproductive and developmental toxicity studies have been conducted recently by this laboratory. All reports published to date have indicated that there are no reproductive, developmental or fetotoxic

effects of this chemical under the test conditions.

Result: Groups of male and female rats were dosed with the test

article at the above levels prior to mating. Males and females from the same dose levels were paired. Animals were observed for body weight, weight gain, food consumption, appearance, behavior, copulation index and fertility index during the life phase of the study. Mated females were

sacrificed on Day 14 of gestation and the fetuses removed via Cesarean Section. Fetuses were weighed, sexed and examined for external, skeletal and soft tissue anomalies as well as

developmental variation

General parental toxicity: All animals survived until planned sacrifice. There were no effects of treatment observed on mean body weight, weight gain, appearance, behavior, physical viability, copulation index or fertility index. There were no remarkable findings in gross necropsy or organ weights.

Toxicity to offspring: The number of corpora lutea and implantations, implantation rate, fetal mortality, and number

of live fetuses were not affected by the test article.

Reliability: (2) valid with restrictions

Meets generally accepted scientific standards, well documented

and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (79)

Type: other: Three generation study

Species: rat Sex: male/female

Strain: other: Charles river CD

Route of admin.: oral feed

Exposure Period: for three successive generations

Frequency of

treatment: daily

Duration of test:

Doses: 100, 300, 1000 ppm (8, 23, 75 mg/kg b.w./day)

Control Group: yes, concurrent no treatment

NOAEL Parental: 10 ppm

Method: other: the F0-generation received the test compound for 11

weeks before mating and during mating, gestation and lactation

for two successive litters (Fla, Flb)

Year: GLP:

Test substance:

Remark: the calculation of the dose levels is based on 1 ppm

corresponds to 0.075 mg/kg b.w.

Result: F0-generation: no effect on fertility, no effect on

behaviour, reduced body weight gain at the mid and high dose

levels, no substance-related histopathological effects F1-generation, F2-generation, F3-generation: no effect on fertility, no effect on behaviour, no substance-related

histopathological effects

Source: Bayer AG Leverkusen

Reliability: (2) valid with restrictions
Flag: Critical study for SIDS endpoint

20-NOV-2001 (80) (52)

- 48/57 -

Date: 20-NOV-2001 5. Toxicity ID: 793-24-8

Type: other: rangefinding study

Species: Sex: female rat

Strain: no data Route of admin.: gavage

Exposure Period: gestation days 6 to 15

Frequency of

treatment: daily

Duration of test:

100, 300, 600, 1000 or 2000 mg/kg bw/d Doses:

Control Group: ves

Method:

GLP: no data Year:

Test substance: other TS: Santoflex 13

excessive toxicity was noted at 600 mg/kg bw/d and

above; intrauterine survival was not affected by

treatment at 100 or 300 mg/kg bw/d

Source: MonsantoBayer AG Leverkusen

31-MAY-1994 (81)

5.9 Developmental Toxicity/Teratogenicity

Sex: female Species: rat

Strain: Sprague-Dawley

Route of admin.: gavage

Exposure period: days 6-15 of gestation

Frequency of

treatment: daily Duration of test: 20 days

0, 50, 100 or 250 mg/kg bw/dDoses: Control Group: yes, concurrent vehicle

NOAEL Maternalt.: = 50 mg/kg bw NOAEL Teratogen.: > 250 mg/kg bw

Method: other: Teratology - Principles and Techniques, J.G. Wilson

1965

Year: GLP: yes

other TS: 6PPD Lot# KE-10-143 purity: >97% Test substance:

Remark: Four groups of 25 bred female rats were dosed with the test

> article at 0, 50, 100 and 250 mg/kg/body weight. Dosages were determined in a preceding range-finding study. Survival was 100% in all groups. Throughout gestation, all animals were observed 2x/day for appearance, behavior, body weight and food consumption. On Day 20, all test animals were sacrificed and

the fetuses removed via Cesarian Section. Fetuses were weighed, sexed and examined for external, skeletal and soft

tissue anomalies as well as developmental variation.

This was a follow-up study to a range-finding study (Monsanto WI-85-304) that noted excessive maternal toxicity at dose levels of 2000, 1000 and 600 mg/kg/day, with clinical signs of toxicity in the 300 mg/kg/day group. Intrauterine survival was not affected at the 100 and 300 mg/kg/day dose levels. Maternal general toxicity: Clinical signs noted in the Mid- to

Result:

High-dose groups included salivation prior to dosing, soft stool, diarrhea and green fecal discoloration. Maternal body weights and weight gain were comparable in all groups. No

morphopathological changes which could be attributed to the test article were observed in any of the treated animals Pregnancy/litter data: No abortions or premature deliveries occurred in any test group.

Foetal data: No differences that could be associated with the test article were observed between the control group and the treated groups with respect to number of viable fetuses, early and late resorptions, fetal sex ratios or fetal weights. The types of malformations and the frequency of such mutations occurring during this study were not those indicative of a teratogenic response. There was a small, non-statistically significant increase in the incidence and number of skeletal variations in the treated groups. However, these were judged to be common developmental variations of this species and have been observed to occur with similar incidence in the historical data.

Not teratogenic or embryo/fetotoxic under test conditions.

Reliability: (1) valid without restriction

GLP study, meets generally accepted scientific standards, well

documented and acceptable for assessment

Flag: Critical study for SIDS endpoint

20-NOV-2001 (82)

Species: rabbit Sex: female

Strain: other: New Zealand Route of admin.: oral unspecified

Exposure period: gestation day 6 through day 18 inclusive

Frequency of

treatment: once a day

Duration of test: post observation: sacrifice on gestation day 29

Doses: 10, 30 mg/kg b.w./day

Control Group: other: yes, empty gelatin capsules

NOAEL Maternalt.: 30 mg/kg bw

Method:

Year: GLP:

Test substance: other TS: Santoflex 13

Remark: in a pilot study 100 and 300 mg/kg b.w./day caused maternal

toxicity

Result: maternal body weight loss and mortality comparable to the

controls, no treatment related gross lesions were noted at necropsy; a slight increase in the number of resorption sites per 100 implantation sites for the 30 mg/kg b.w. group (38.6 %) when compared to the controls (31.4 %), the number of live young per 100 implantation sites for the 10 mg/kg b.w. group (48.3 %) and for the 30 mg/kg b.w. group (38.6 %) were moderately decreased when compared to the controls (68.6 %); no increase in the incidence of external, visceral

and skeletal abnormalities

Source: Bayer AG Leverkusen

20-NOV-2001 (83)

- 50/57 -

Date: 20-NOV-2001
5. Toxicity ID: 793-24-8

Species: Sex:

Strain:

Route of admin.:
Exposure period:
Frequency of
 treatment:
Duration of test:

Doses:

Control Group:

Method: other: test compounds were tested for embryotoxicity and induction of malformations in three-day chicken embryos

Year: GLP:

Test substance:

Result: slight effects

Source: Bayer AG Leverkusen

08-DEC-1992 (84) (85)

5.10 Other Relevant Information

Type: other

Remark: A comprehensive description of the toxicity profile is

available in the BUA-Report

Source: Bayer AG Leverkusen

12-NOV-1998 (86)

Type:

Remark: Revision date: August, 1998

Source: Bayer AG Leverkusen

17-AUG-1998

5.11 Experience with Human Exposure

Memo: Occupational eczema study - 6PPD and IPPD exposures

Remark: Cross sensitization in rubber workers exposed to various

members of the PPD family have been reported. Anecdotal evidence suggests that this class of compounds has a high potential for skin sensitization with prolonged and repeated

exposures of sensitive individuals.

20-NOV-2001 (87)

Remark: In the rubber industry 6PPD was detected in the urine of

6PPD exposed workers

Source: Bayer AG Leverkusen

08-DEC-1992 (88)

Remark: analytical methods for the determination of the trace levels

of 6PPD in human urine are described (in the publication of Pavan et. al the abbreviation 6PPD is used however the substance is called N-(2,3-dimethylpropyl)-N-phenyl-

1,4-benzenediamine with the CAS-No. 739-24-8)

Source: Bayer AG Leverkusen

08-DEC-1992 (89) (90)

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- (2) Hawley, G.G.: The Condensed Chemical Dictionary, 9th ed., 672 (1977)
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- (8) Tests Monsanto, cited in Monsanto Material Safety Data Sheet Santoflex 13, Antioxonant, 4186
- (9) Monsanto ES-78-SS-20 MIC Environmental Science Dec. 1978
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- (11) Meylan W. and Howard P. (1999) EPIWin Modeling Program. Syracuse Research Corporation. Environmental Science Center, 6225 Running Ridge Road, North Syracuse, NY 13212-2510.
- (12) Monsanto #32304. Analytical BioChemistry Labs. March, 1986
- (13) Monsanto study ES-78-SS28
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- (16) Monsanto ES-81-SS-52 Environmental Sciences Labs Dec. 1981
- (17) Study Bayer AG, WV-LE Umweltschutz, AWALU, 1984
- (18) Simpson, K.E., Int. Biodeterioration 24, 307-312 (1988)
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- (21) MonsantoAB-78-121-B Analytical BioChemistry Labs July 1979
- (22) Monsanto AB-78-121 Analytical BioChemistry Labs, June 1978
- (23) Monsanto Study MO-92-9043 (ES-81-SS-36)

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- (25) Monsanto report MSL-6158
- (26) Monsanto BN-78-362 EG&G Bionomics Sept. 1978
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- (30) Stasenkova, K.P., Sov. Rubber Technol. 29, 25-26 (1970)
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- (32) Goodyear, August 8, 1973
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- (34) Monsanto SH-76-8 Industrial Biology Laboratories 1976
- (35) American Cyanamid Company data (1972), NTIS/OTS 206438, EPA/OTS document 878213691 (1983)
- (36) Monsanto data, Project No. SH-64-8, Experiment No. 30-11, Stocks for Dermatitis Studies, June 11, 1964
- (37) Monsanto data, Project No. SH-64-5, March 11, 1964
- (38) Monsanto data, Project No. SH-64-6, Experiment No. 16-91, Comparison of n-Hexyl Analog with Santoflex 13, March 23, 1964
- (39) Monsanto data, Project No. SH-63-15, Experiment No. 16-64, Stocks for Dermatitis Studies, November 15, 1963
- (40) Monsanto data, Project No. SH-63-5, Experiment No. 11-55, Composition of Rubber, Stocks for Dermatitis Studies, April 30, 1963
- (41) Monsanto data, Project No. SH-63-14, November 8, 1963
- (42) Monsanto data, Project No. SH-64-4, March 6, 1964
- (43) Monsanto data, Dermatitic effects of Santoflex IP and Santoflex 13

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- (51) Monsanto BTL-74-26 Industrial Bio-Test Labs Nov. 27, 1978.
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7.1 End Point Summary

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7.2 Hazard Summary

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7.3 Risk Assessment

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